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- (71) Applicant: TRIMERIS, INC. [US/US]; 4727 University Drive, Durham, NC 27707 (US).
- (72) Inventors: JEFFS, Peter; 200 Redbud Lane, Chapel Hill, NC 27514 (US). LACKEY, John, William; 3220 Wood Duck Lane, Hillsborough, NC 27278 (US). ERICKSON, Joel, Burton; 5408 Beumont Drive, Durham, NC 27707 (US). LAWLESS, Mary, K.; 10521 New Arden Way, Raleigh, NC 27613 (US). MERUTKA, Gene; 18456 Purdue Drive, Saratoga, CA 95070 (US).
- (74) Agents: CORUZZI, Laura, A. et al.; Pennie & Edmonds LLP, 1155 Avenue of the Americas, New York, NY 10036 (US).

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(54) Title: METHODS AND COMPOSITIONS FOR INHIBITION OF MEMBRANE FUSION-ASSOCIATED EVENTS, IN-CLUDING HIV TRANSMISSION

(57) Abstract: The present invention relates to peptides which exhibit potent anti-retroviral activity. The peptides of the invention comprise DP178 (SEQ ID:1) peptide corresponding to amino acids 638 to 673 of the HIV-1_{LAI} gp41 protein, and fragments, analogs and homologs of DP178. The invention further relates to the uses of such peptides as inhibitory of human and non-human retroviral, especially HIV, transmission to uninfected cells.

METHODS AND COMPOSITIONS FOR INHIBITION OF MEMBRANE FUSION-ASSOCIATED EVENTS, INCLUDING HIV TRANSMISSION

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1. INTRODUCTION

The present invention relates, first, to DP178 (SEQ ID NO:1), a peptide, also referred to herein as T20, corresponding to amino acids 638 to 673 of the HIV-1LMI transmembrane protein (TM) gp41, and portions or analogs of DP178 (SEQ ID NO:1), which exhibit antimembrane fusion capability, antiviral activity, such as the ability to inhibit HIV transmission to uninfected CD-4° cells, or an ability to modulate intracellular processes involving coiled-coil peptide structures. The present invention also relates to peptides analogous to DP107 (SEQ ID NO:25), a peptide, which is also referred to herein as T21, corresponding to amino acids 558 to 595 of the HIV-1, transmembrane protein (TM) gp41, having amino acid sequences present 20 in other viruses, such as enveloped viruses, and/or other organisms, and further relates to the uses of such peptides. These peptides exhibit anti-membrane fusion capability, antiviral activity, or the ability to modulate intracellular processes involving coiledcoil peptide structures.

The gp41 region from which DP107 is derived is referred to herein as HR1. The gp41 region from which DP178 is derived is referred to herein as HR2. As discussed herein, the gp41 HR1 and HR2 regions interact (non-covalently) with each other and/or with

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T20 and T21 peptides. This interaction is required for normal infectivity of HIV.

The present invention therefore additionally relates to methods for identifying compounds, including small molecule compounds, that disrupt the 5 interaction between DP178 and DP107, and/or between DP107-like and DP178-like peptides. In one embodiment, such methods relate to identification and utilization of modified DP178, DP178-like, DP107 and DP107-like peptides and peptide pairs that interact 10 with each other at a lower affinity than the affinity exhibited by corresponding "parent" or "native" peptides. Further, the invention relates to the use of DP178, DP178 portions, DP107, DP017 portions and/or analogs and other modulators, including small molecules modulators, of DP178/DP107, 15 DP178-like/DP107-like, or HR1/HR2 interactions as antifusogenic or antiviral compounds or as inhibitors of intracellular events involving coiled-coil peptide structures. The invention is demonstrated, first, by way of an Example wherein DP178 (SEQ ID:1), and a peptide whose sequence is homologous to DP178 are each shown to be potent, non-cytotoxic inhibitors of HIV-1 transfer to uninfected CD-4 cells. The invention is further demonstrated by Examples wherein peptides having structural and/or amino acid motif similarity 25 to DP107 and DP178 are identified in a variety of viral and nonviral organisms, and in examples wherein a number of such identified peptides derived from several different viral systems are demonstrated to exhibit antiviral activity. The invention is still further demonstrated by way of Examples wherein other 30

DP178-like and DP107-like peptides are identified that

interact with their corresponding HR1 and HR2 domains with a lower affinity than the affinity exhibited by the native DP178 or DP107 peptide from which they are derived.

2. BACKGROUND OF THE INVENTION

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2.1 MEMBRANE FUSION EVENTS

Membrane fusion is a ubiquitous cell biological process (for a review, see White, J.M., 1992, Science 258:917-924). Fusion events which mediate cellular housekeeping functions, such as endocytosis, constitutive secretion, and recycling of membrane components, occur continuously in all eukaryotic cells.

Additional fusion events occur in specialized

cells. Intracellularly, for example, fusion events

are involved in such processes as occur in regulated
exocytosis of hormones, enzymes and neurotransmitters.
Intercellularly, such fusion events feature
prominently in, for example, sperm-egg fusion and
myoblast fusion.

Fusion events are also associated with disease states. For example, fusion events are involved in the formation of giant cells during inflammatory reactions, the entry of all enveloped viruses into cells, and, in the case of human immunodeficiency virus (HIV), for example, are responsible for the virally induced cell-cell fusion which leads to cell death.

2.2. THE HUMAN IMMUNODEFICIENCY VIRUS

The human immunodeficiency virus (HIV) has been implicated as the primary cause of the slowly

degenerative immune system disease termed acquired immune deficiency syndrome (AIDS) (Barre-Sinoussi, F. et al., 1983, Science 220:868-870; Gallo, R. et al., 1984, Science 224:500-503). There are at least two distinct types of HIV: HIV-1 (Barre-Sinoussi, F. et al., 1983, Science 220:868-870; Gallo R. et al., 1984, Science 224:500-503) and HIV-2 (Clavel, F. et al., 1986, Science 233:343-346; Guyader, M. et al., 1987, Nature 326:662-669). Further, a large amount of genetic heterogeneity exists within populations of each of these types. Infection of human CD-4* T-lymphocytes with an HIV virus leads to depletion of the cell type and eventually to opportunistic infections, neurological dysfunctions, neoplastic growth, and ultimately death.

HIV is a member of the lentivirus family of retroviruses (Teich, N. et al., 1984, RNA Tumor Viruses, Weiss, R. et al., eds., CSH-Press, pp. 949-956). Retroviruses are small enveloped viruses that contain a diploid, single-stranded RNA genome, and replicate via a DNA intermediate produced by a virally-encoded reverse transcriptase, an RNA-dependent DNA polymerase (Varmus, H., 1988, Science 240:1427-1439). Other retroviruses include, for example, oncogenic viruses such as human T-cell leukemia viruses (HTLV-I,-II,-III), and feline leukemia virus.

The HIV viral particle consists of a viral core, composed of capsid proteins, that contains the viral RNA genome and those enzymes required for early replicative events. Myristylated Gag protein forms an outer viral shell around the viral core, which is, in turn, surrounded by a lipid membrane enveloped derived

from the infected cell membrane. The HIV enveloped surface glycoproteins are synthesized as a single 160 Kd precursor protein which is cleaved by a cellular protease during viral budding into two glycoproteins, gp41 and gp120. gp41 is a transmembrane protein and gp120 is an extracellular protein which remains noncovalently associated with gp41, possibly in a trimeric or multimeric form (Hammarskjold, M. and Rekosh, D., 1989, Biochem. Biophys. Acta 989:269-280).

HIV is targeted to CD-4 cells because the CD-4 cell surface protein acts as the cellular receptor for the HIV-1 virus (Dalgleish, A. et al., 1984, Nature 312:763-767; Klatzmann et al., 1984, Nature 312:767-768; Maddon et al., 1986, Cell 47:333-348). Viral entry into cells is dependent upon gp120 binding the cellular CD-4 receptor molecules (McDougal, J.S. et al., 1986, Science 231:382-385; Maddon, P.J. et al., 1986, Cell 47:333-348) and thus explains HIV's tropism for CD-4+ cells, while gp41 anchors the enveloped glycoprotein complex in the viral membrane.

20 2.3. HIV TREATMENT

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HIV infection is pandemic and HIV associated diseases represent a major world health problem. Although considerable effort is being put into the successful design of effective therapeutics, currently 25 no curative anti-retroviral drugs against AIDS exist. In attempts to develop such drugs, several stages of the HIV life cycle have been considered as targets for therapeutic intervention (Mitsuya, H. et al., 1991, FASEB J. 5:2369-2381). For example, virally encoded reverse transcriptase has been one focus of drug development. A number of reverse-transcriptase-

targeted drugs, including 2',3'-dideoxynucleoside
analogs such as AZT, ddI, ddC, and d4T have been
developed which have been shown to been active against
HIV (Mitsuya, H. et al., 1991, Science 249:1533-1544).
While beneficial, these nucleoside analogs are not
curative, probably due to the rapid appearance of drug
resistant HIV mutants (Lander, B. et al., 1989,
Science 243:1731-1734). In addition, the drugs often
exhibit toxic side effects such as bone marrow
suppression, vomiting, and liver function
abnormalities.

which can inhibit viral entry into the cell, the earliest stage of HIV infection. Here, the focus has thus far been on CD4, the cell surface receptor for HIV. Recombinant soluble CD4; for example, has been shown to inhibit infection of CD-4* T-cells by some HIV-1 strains (Smith, D.H. et al., 1987, Science 238:1704-1707). Certain primary HIV-1 isolates, however, are relatively less sensitive to inhibition by recombinant CD-4 (Daar, E. et al., 1990, Proc. Natl. Acad. Sci. USA 87:6574-6579). In addition,

Natl. Acad. Sci. USA <u>87</u>:6574-6579). In addition, recombinant soluble CD-4 clinical trials have produced inconclusive results (Schooley, R. <u>et al.</u>, 1990, Ann. Int. Med. <u>112</u>:247-253; Kahn, J.O. <u>et al.</u>, 1990, Ann. Int. Med. <u>112</u>:254-261; Yarchoan, R. <u>et al.</u>, 1989, Proc. Vth Int. Conf. on AIDS, p. 564, MCP 137).

The late stages of HIV replication, which involve crucial virus-specific secondary processing of certain viral proteins, have also been suggested as possible anti-HIV drug targets. Late stage processing is dependent on the activity of a viral protease, and drugs are being developed which inhibit this protease

(Erickson, J., 1990, Science <u>249</u>:527-533). The clinical outcome of these candidate drugs is still in question.

Attention is also being given to the development of vaccines for the treatment of HIV infection. 5 HIV-1 enveloped proteins (gp160, gp120, gp41) have been shown to be the major antigens for anti-HIV antibodies present in AIDS patients (Barin, et al., 1985, Science 228:1094-1096). Thus far, therefore, these proteins seem to be the most promising 10 candidates to act as antigens for anti-HIV vaccine development. To this end, several groups have begun to use various portions of gp160, gp120, and/or gp41 as immunogenic targets for the host immune system. See for example, Ivanoff, L. et al., U.S. Pat. No. 5,141,867; Saith, G. et al., WO 92/22,654; Shafferman, 15 A., WO 91/09,872; Formoso, C. et al., WO 90/07,119. Clinical results concerning these candidate vaccines, however, still remain far in the future.

Thus, although a great deal of effort is being directed to the design and testing of anti-retroviral drugs, a truly effective, non-toxic treatment is still needed.

3. SUMMARY OF THE INVENTION

The present invention relates, first, to DP178, a
36-amino acid synthetic peptide, also referred to
herein as T20, corresponding to amino acids 638 to 673
of the transmembrane protein (TM) gp41 from the HIV-1
isolate LAI (HIV-1_{LAI}), which exhibits potent antiHIV-1 activity. The gp41 region from which DP178 is
derived in referred to herein as HR2.

The invention further relates to those portions and analogs of DP178 which also show such antiviral activity, and/or show anti-membrane fusion capability, or an ability to modulate intracellular processes involving coiled-coil peptide structures. The term 5 "DP178 analog" refers to a peptide which contains an amino acid sequence corresponding to the DP178 peptide sequence present within the gp41 protein of HIV-1, , but found in viruses and/or organisms other than HIV-1 LAI. Such DP178 analog peptides may, therefore, 10 correspond to DP178-like amino acid sequences present in other viruses, such as, for example, enveloped viruses, such as retroviruses other than HIV-1 LAI, as well as non-enveloped viruses. Further, such analogous DP178 peptides may also correspond to DP178like amino acid sequences present in nonviral organisms.

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The invention further relates to DP107, a peptide, which is also referred to herein as T21, corresponding to amino acids 558-595 of the HIV-1, at transmembrane protein (TM) gp41. The gp41 region from 20 which DP107 is derived is referred to herein as HR1. The invention also relates to those portions and analogs of DP107 which that also show antiviral activity, and/or show anti-membrane fusion capability, or an ability to modulate intracellular processes 25 involving coiled-coil peptide structures. The term "DP107 analog" as used herein refers to a peptide which contains an amino acid sequence corresponding to the DP107 peptide sequence present within the gp41 protein of HIV-1LAI, but found in viruses and organisms other than HIV-1_{LAI}. Such DP107 analog peptides may, 30 therefore, correspond to DP107-like amino acid

sequences present in other viruses, such as, for for example, enveloped viruses, such as retroviruses other than HIV-1_{LAI}, as well as non-enveloped viruses. Further, such DP107 analog peptides may also correspond to DP107-like amino acid sequences present in nonviral organisms.

Further, the peptides of the invention include DP107 analog and DP178 analog peptides having amino acid sequences recognized or identified by the 107x178x4, ALLMOTI5 and/or PLZIP search motifs described herein.

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The peptides of the invention may, for example, exhibit antifusogenic activity, antiviral activity, and/or may have the ability to modulate intracellular processes which involve coiled-coil peptide structures. With respect to the antiviral activity of 15 the peptides of the invention, such an antiviral activity includes, but is not limited to the inhibition of HIV transmission to uninfected CD-4* cells. Additionally, the antifusogenic capability, antiviral activity or intracellular modulatory activity of the peptides of the invention merely requires the presence of the peptides of the invention, and, specifically, does not require the stimulation of a host immune response directed against such peptides.

25 The peptides of the invention may be used, for example, as inhibitors of membrane fusion-asociated events, such as, for example, the inhibition of human and non-human retroviral, especially HIV, transmission to uninfected cells. It is further contemplated that the peptides of the invention may be used as

modulators of intracellular events involving coiledcoil peptide structures.

The peptides of the invention may, alternatively, be used to identify compounds, including small molecule compounds, which may themselves exhibit antifusogenic, antiviral, or intracellular modulatory activity. For example, in one embodiment, the peptides of the invention are used to identify other DP178-like and/or DP107-like peptides that interact with each other and/or with their complementary HR1 or 10 HR2 domains with a lower affinity than the affinity exhibited by the "parent" or "native" DP178 or DP107 peptides from which they are derived. Such DP178-like and DP107-like peptides, which are also part of the present invention, may also be used, e.g., to identify compounds, such as small molecule compounds, that 15 exhibit antifusogenic, antiviral, or intracellular modulatory activity.

Additional uses include, for example, the use of the peptides of the invention as organism or viral type and/or subtype-specific diagnostic tools.

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The terms "antifusogenic" and "anti-membrane fusion", as used herein, refer to an agent's ability to inhibit or reduce the level of membrane fusion events between two or more moieties relative to the level of membrane fusion which occurs between said 25 moieties in the absence of the peptide. The moieties may be, for example, cell membranes or viral structures, such as viral envelopes or pili. "antiviral", as used herein, refers to the compound's ability to inhibit viral infection of cells, via, for example, cell-cell fusion or free virus infection. Such infection may involve membrane fusion, as occurs

in the case of enveloped viruses, or some other fusion event involving a viral structure and a cellular structure (e.g., such as the fusion of a viral pilus and bacterial membrane during bacterial conjugation).

It is also contemplated that the peptides of the invention may exhibit the ability to modulate intracellular events involving coiled-coil peptide structures. "Modulate", as used herein, refers to a stimulatory or inhibitory effect on the intracellular process of interest relative to the level or activity 10 of such a process in the absence of a peptide of the invention.

Embodiments of the invention are demonstrated below wherein an extremely low concentration of DP178 (SEQ ID:1), and very low concentrations of a DP178 homolog (SEQ ID:3) are shown to be potent inhibitors of HIV-1 mediated CD-4 cell-cell fusion (i.e., syncytial formation) and infection of CD-4 cells by cell-free virus. Further, it is shown that DP178 (SEQ ID:1) is not toxic to cells, even at concentrations 3 logs higher than the inhibitory DP-178 (SEQ ID:1) concentration.

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The present invention is based, in part, on the surprising discovery that the DP107 and DP178 domains of the HIV gp41 protein non-covalently complex with each other, and that their interaction is required for 25 the normal infectivity of the virus. This discovery is described in the Example presented, below, in Section 8. The invention, therefore, further relates to methods for identifying antifusogenic, including antiviral, compounds that disrupt the interaction between DP107 and DP178, and/or between DP107-like and DP178-like peptides.

Additional embodiments of the invention

(specifically, the Examples presents in Sections 9-16 and 19-25, below) are demonstrated, below, wherein peptides, from a variety of viral and nonviral sources, having structural and/or amino acid motif

5 similarity to DP107 and DP178 are identified, and search motifs for their identification are described. Further, Examples (in Sections 17, 18, 25-29) are presented wherein a number of the peptides of the invention are demonstrated exhibit substantial

10 antiviral activity or activity predictive of antiviral activity.

3.1. DEFINITIONS

comprising two or more amino acids covalently joined by peptide bonds. Peptides may be referred to with respect to the number of constituent amino acids, i.e., a dipeptide contains two amino acid residues, a tripeptide contains three, etc. Peptides containing ten or fewer amino acids may be referred to as oligopeptides, while those with more than ten amino acid residues are polypeptides. Such peptides may also include any of the modifications and additional amino and carboxy groups as are described herein.

Peptide sequences defined herein are represented 25 by one-letter symbols for amino acid residues as follows:

- A (alanine)
- R (arginine)
- N (asparagine)
- D (aspartic acid)
- C (cysteine)
- 30 Q (glutamine)
 - E (glutamic acid)

G (glycine) H (histidine) I (isoleucine) L (leucine) K (lysine) M (methionine) F (phenylalanine) 5 P (proline) S (serine) T (threonine) W (tryptophan) Y (tyrosine) V (valine)

4. BRIEF DESCRIPTION OF THE FIGURES

10 FIG. 1. Amino acid sequence of DP178 (SEQ ID:1) derived from HIV, DP178 homologs derived from HIV-1, FP2 (DP-185; SEQ ID:3), HIV-1_{RF} (SEQ ID:4), and HIV-1_{MN} (SEQ ID:5); DP178 homologs derived from amino acid sequences of two prototypic HIV-2 isolates, namely, 15 HIV-2_{rod} (SEQ ID:6) and HIV-2_{NIHZ} (SEQ ID:7); control peptides: DP-180 (SEQ ID:2), a peptide incorporating the amino acid residues of DP178 in a scrambled sequence; DP-118 (SEQ ID:10) unrelated to DP178, which inhibits HIV-1 cell free virus infection; DP-125 (SEQ

20 ID:8), unrelated to DP178, also inhibits HIV-1 cell free virus infection; DP-116 (SEQ ID:9), unrelated to DP178, is negative for inhibition of HIV-1 infection when tested using a cell-free virus infection assay. Throughout the figures, the one letter amino acid code is used.

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Inhibition of HIV-1 cell-free virus infection by synthetic peptides. ICso refers to the concentration of peptide that inhibits RT production from infected cells by 50% compared to the untreated control. Control: the level of RT produced by

untreated cell cultures infected with the same level of virus as treated cultures.

FIG. 3. Inhibition of HIV-1 and HIV-2 cell-free virus infection by the synthetic peptide DP178 (SEQ ID:1). IC₅₀: concentration of peptide that inhibits RT production by 50% compared to the untreated control. Control: Level of RT produced by untreated cell cultures infected with the same level of virus as treated cultures.

pp178 (SEQ ID:1) inhibition of HIV-1 prototypic isolate-mediated syncytial formation; data represents the number of virus-induced syncytial per cell. FIG. 4B: DP-180 (SEQ ID:2) represents a scrambled control peptide; DP-185 (SEQ ID:3) represents a DP178 homolog derived from HIV-1_{SF2} isolate; Control, refers to the number of syncytial produced in the absence of peptide.

FIG. 5. Fusion inhibition assay: HIV-1 vs. HIV-2. Data represents the number of virus-induced syncytial per well. ND: not done.

FIG. 6. Cytotoxicity study of DP178 (SEQ ID:1) and DP-116 (SEQ ID:9) on CEM cells. Cell proliferation data is shown.

FIG. 7. Schematic representation of HIV-gp41 and maltose binding protein (MBP)-gp41 fusion

proteins. DP107 and DP178 are synthetic peptides based on the two putative helices of gp41. The letter P in the DP107 boxes denotes an Ile to Pro mutation at amino acid number 578. Amino acid residues are numbered according to Meyers et al., "Human

Retroviruses and AIDS", 1991, Theoret. Biol. and
Biophys. Group, Los Alamos Natl. Lab., Los Alamos, NM.

The proteins are more fully described, below, in Section 8.1.1.

- FIG. 8. A point mutation alters the conformation and anti-HIV activity of M41.
- FIG. 9. Abrogation of DP178 anti-HIV activity.

 5 Cell fusion assays were carried out in the presence of
 10 nM DP178 and various concentrations of M41Δ178 or
 M41PΔ178.
 - FIG. 10. Binding of DP178 to leucine zipper of qp41 analyzed by FAb-D ELISA.
- Models for a structural transition FIG. 11A-B. 10 in the HIV-1 TM protein. Two models are proposed which indicate a structural transition from a native oligomer to a fusogenic state following a trigger event (possibly gp120 binding to CD4). Common features of both models include (1) the native state 15 is held together by noncovalent protein-protein * interactions to form the heterodimer of qp120/41 and other interactions, principally though gp41 interactive sites, to form homo-oligomers on the virus surface of the gp120/41 complexes; (2) shielding of 20 the hydrophobic fusogenic peptide at the N-terminus (F) in the native state; and (3) the leucine zipper domain (DP107) exists as a homo-oligomer coiled coil only in the fusogenic state. The major differences in the two models include the structural state (native or 25 fusogenic) in which the DP107 and DP178 domains are
- fusogenic) in which the DP107 and DP178 domains are complexed to each other. In the first model (FIG. 11A) this interaction occurs in the native state and in the second (FIG. 11B), it occurs during the fusogenic state. When triggered, the fusion complex in the model depicted in (A) is generated through

DP107 domains resulting in an extended α-helix. This conformational change positions the fusion peptide for interaction with the cell membrane. In the second model (FIG. 11B), the fusogenic complex is stabilized by the association of the DP178 domain with the DP107 coiled-coil.

- FIG. 12. Motif design using heptad repeat positioning of amino acids of known coiled-coils.
- FIG. 13. Motif design using proposed heptad repeat positioning of amino acids of DP107 and DP178.
- 10 FIG. 14. Hybrid motif design crossing GCN4 and DP107.
 - FIG. 15. Hybrid motif design crossing GCN4 and DP178.
- FIG. 16. Hybrid motif design 107x178x4,

 crossing DP107 and DP178. This motif was found to be
 the most consistent at identifying relevant DP107-like
 and DP178-like peptide regions.
 - FIG. 17. Hybrid motif design crossing GCN4, DP107, and DP178.
- FIG. 18. Hybrid motif design ALLMOTI5

 crossing GCN4, DP107, DP178, c-Fos c-Jun, c-Myc, and
 Flu Loop 36.
 - FIG. 19. PLZIP motifs designed to identify N-terminal proline-leucine zipper motifs.
- FIG. 20. Search results for HIV-1 (BRU

 25 isolate) enveloped protein gp41. Sequence search
 motif designations: Spades (♠): 107x178x4; Hearts (♥)

 ALLMOTI5; Clubs (♠): PLZIP; Diamonds (♠):
 transmembrane region (the putative transmembrane
 domains were identified using a PC/Gene program
 designed to search for such peptide regions).
 - Asterisk (*): Lupas method. The amino acid sequences

identified by each motif are bracketed by the respective characters. Representative sequences chosen based on 107x178x4 searches are underlined and in bold. DP107 and DP178 sequences are marked, and additionally double-underlined and italicized.

- FIG. 21. Search results for human respiratory syncytial virus (RSV) strain A2 fusion glycoprotein F1. Sequence search motif designations are as in FIG. 20.
- FIG. 22. Search results for simian

 immunodeficiency virus (SIV) enveloped protein gp41

 (AGM3 isolate). Sequence search motif designations
 are as in FIG. 20.
- FIG. 23. Search results for canine
 distemper virus (strain Onderstepoort) fusion
 glycoprotein 1. Sequence search motif designations
 are as in FIG. 20.
 - FIG. 24. Search results for newcastle disease virus (strain Australia-Victoria/32) fusion glycoprotein F1. Sequence search motif designations are as in FIG. 20.
- FIG. 25. Search results for human parainfluenza 3 virus (strain NIH 47885) fusion glycoprotein F1. Sequence search motif designations are as in FIG. 20.
- FIG. 26. Search results for influenza A
 25 virus (strain A/AICHI/2/68) hemagglutinin precursor
 HA2. Sequence search designations are as in FIG. 20.

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FIG. 27A-D. Respiratory Syncytial Virus (RSV) peptide antiviral and circular dichroism data.
FIG. 27A-B: Peptides derived from the F2 DP178/DP107-like region. Antiviral and CD data. FIG. 27C-D:

Peptides derived from the F1 DP107-like region. Peptide and CD data.

Antiviral activity (AV) is represented by the following qualitative symbols:

"-", negative antiviral activity;

5 "+/-", antiviral activity at greater than $100\mu g/ml$;

"+", antiviral activity at between $50-100\mu g/ml$;

"++", antiviral activity at between $20-50\mu g/ml$;

"+++", antiviral activity at between 1-20 μ g/ml;

"++++", antiviral activity at $<1\mu$ g/ml.

CD data, referring to the level of helicity is represented by the following qualitative symbol:

"-", no helicity;

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"+", 25-50% helicity;

"++", 50-75% helicity;

"+++"' 75-100% helicity.

 IC_{50} refers to the concentration of peptide necessary to produce only 50% of the number of syncytial relative to infected control cultures containing no peptide. IC_{50} values were obtained using purified peptides only.

FIG. 28A-B. Respiratory Syncytial Virus (RSV) DP178-like region (F1) peptide antiviral and CD data. Antiviral symbols, CD symbols, and IC₅₀ are as in FIG. 27A-D. IC₅₀ values were obtained using purified peptides only.

FIG. 29A-B. Peptides derived from the HPIV3 F1 DP107-like region. Peptide antiviral and CD data. Antiviral symbols, CD symbols, and IC_{50} are as in FIG. 27A-D. Purified peptides were used to obtain IC_{50} values, except where the values are marked by an

asterisk (*), in which cases, the IC_{50} values were obtained using a crude peptide preparation.

FIG. 29C. HPIV3 peptide T-184 CD spectrum at 1°C in 0.1M NaCl 10mM KPO₄, pH 7.0. The data demonstrates the peptide's helical secondary structure $(\theta_{222/208}=1.2)$ over a wide range of concentrations (100-1500 μ M). This evidence is consistent with the peptide forming a helical coiled-coil structure.

FIG. 30A-B. Peptides derived from the HPIV3 F1 DP178-like region. Peptide antiviral and CD data.

- Antiviral symbols, CD symbols, and IC_{50} are as in FIG. 27A-D. Purified peptides were used to obtain IC_{50} values, except where the values are marked by an asterisk (*), in which cases, the IC_{50} values were obtained using a crude peptide preparation.
- FIG. 31. Motif search results for simian immunodeficiency virus (SIV) isolate MM251, enveloped polyprotein gp41. Sequence search designations are as in FIG. 20.

FIG. 32. Motif search results for Epstein-Barr Virus (Strain B95-8), glycoprotein gpll0
precursor (designated gpll5). BALF4. Sequence search designations are as in FIG. 20.

FIG. 33. Motif search results for Epstein-Barr Virus (Strain B95-8), BZLF1 trans-activator protein (designated EB1 or Zebra). Sequence search designations are as in FIG. 20. Additionally, "@" refers to a well known DNA binding domain and "+" refers to a well known dimerization domain, as defined by Flemington and Speck (Flemington, E. and Speck, S.H., 1990, Proc. Natl. Acad. Sci. USA 87:9459-9463).

FIG. 34. Motif search results for measles

virus (strain Edmonston), fusion glycoprotein F1. Sequence search designations are as in FIG. 20.

FIG. 35. Motif search results for Hepatitis
B Virus (Subtype AYW), major surface antigen precursor
S. Sequence search designations are as in FIG. 20.

FIG. 36. Motif search results for simian Mason-Pfizer monkey virus, enveloped (TM) protein gp20. Sequence search designations are as in FIG. 20.

FIG. 37. Motif search results for Pseudomonas aerginosa, fimbrial protein (Pilin).

10 Sequence search designations are as in FIG. 20.

FIG. 38. Motif search results for Neisseria gonorrhoeae fimbrial protein (Pilin). Sequence search designations are as in FIG. 20.

FIG. 39. Motif search results for

Hemophilus influenzae fimbrial protein. Sequence search designations are as in FIG. 20.

FIG. 40. Motif search results for Staphylococcus aureus, toxic shock syndrome toxin-1. Sequence search designations are as in FIG. 20.

FIG. 41. Motif search results for

Staphylococcus aureus enterotoxin Type E. Sequence search designations are as in FIG. 20.

FIG. 42. Motif search results for Staphylococcus aureus enterotoxin A. Sequence search designations are as in FIG. 20.

25 FIG. 43. Motif search results for Escherichia coli, heat labile enterotoxin A. Sequence search designations are as in FIG. 20.

FIG. 44. Motif search results for human cfos proto-oncoprotein. Sequence search designations are as in FIG. 20.

FIG. 45. Motif search results for human lupus KU autoantigen protein P70. Sequence search designations are as in FIG. 20.

FIG. 46. Motif search results for human zinc finger protein 10. Sequence search designations are as in FIG. 20.

FIG. 47. Measles virus (MeV) fusion protein DP178-like region antiviral and CD data. Antiviral symbols, CD symbols, and IC_{50} are as in FIG. 27A-D. IC_{so} values were obtained using purified peptides.

FIG. 48. Simian immunodeficiency virus 10 (SIV) TM (fusion) protein DP178-like region antiviral data. Antiviral symbols are as in FIG. 27A-D "NT", not tested.

FIG. 49A-C. DP178-derived peptide antiviral The peptides listed herein were derived from 15 the region surrounding the HIV-1 BRU isolate DP178 region (e.g., gp41 amino acid residues 615-717).

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In instances where peptides contained DP178 point mutations, the mutated amino acid residues are shown with a shaded background. In instances in which the test peptide has had an amino and/or carboxy-terminal group added or removed (apart from the standard amidoand acetyl- blocking groups found on such peptides), such modifications are indicated. FIG. 49A: The column to the immediate right of the name of the test 25 peptide indicates the size of the test peptide and points out whether the peptide is derived from a one amino acid peptide "walk" across the DP178 region. The next column to the right indicates whether the test peptide contains a point mutation, while the column to its right indicates whether certain amino acid residues have been added to or removed from the

DP178-derived amino acid sequence. FIG 49B: The column to the immediate right of the test peptide name indicates whether the peptide represents a DP178 truncation, the next column to the right points out whether the peptide contains a point mutation, and the 5 column to its right indicates whether the peptide contains amino acids which have been added to or removed from the DP178 sequence itself. FIG. 49C: The column to the immediate right of the test peptide name indicates whether the test peptide contains a 10 point mutation, while the column to its right indicates whether amino acid residues have been added to or removed from the DP178 sequence itself. IC, is as defined in FIG. 27A-D, and IC₅₀ values were obtained using purified peptides except where marked with an asterisk (*), in which case the IC₅₀ was obtained using 15 a crude peptide preparation.

FIG. 50. DP107 and DP107 gp41 region truncated peptide antiviral data. IC_{50} as defined in FIG. 27A-D, and IC_{50} values were obtained using purified peptides except where marked with an asterisk (*), in which case the IC_{50} was obtained using a crude peptide preparation.

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FIG. 51A-B. Epstein-Barr virus Strain B95-8
BZLF1 DP178/DP107 analog region peptide walks and
electrophoretic mobility shift assay results. The
25 peptides (T-423 to T-446, FIG. 51A; T-447 to T-461,
FIG. 51B) represent one amino acid residue "walks"
through the EBV Zebra protein region from amino acid
residue 173 to 246.

The amino acid residue within this region which corresponds to the first amino acid residue of each peptide is listed to the left of each peptide, while

the amino acid residue within this region which corresponds to the last amino acid residue of each peptide is listed to the right of each peptide. The length of each test peptide is listed at the far right of each line, under the heading "Res".

"ACT" refers to a test peptide's ability to inhibit Zebra binding to its response element. "+" refers to a visible, but incomplete, abrogation of the response element/Zebra homodimer complex; "+++" refers to a complete abrogation of the complex; and "-" represents a lack of complex disruption.

FIG. 52A-B. Hepatitis B virus subtype AYW major surface antigen precursor S protein DP178/DP107 analog region and peptide walks. 52A depicts Domain I (S protein amino acid residues 174-220), which contains a potential DP178/DP107 analog region. In addition, peptides are listed which represent one amino acid peptide "walks" through domain I. 52B depicts Domain II (S protein amino acid residues 233-291), which contains a second potential DP178/DP107 analog region. In addition, peptides are listed which represent one amino acid peptide "walks" through domain II.

FIG. 53: Cell fusion and competitive inhibition data for alanine walk experiments for the DP178-like Respiratory Syncytial Virus (RSV) peptide T112.

FIG. 54: Circular dichroism, cell fusion and competitive inhibition data for alanine walk experiments for the peptide T20, which is also known as DP178.

5. DETAILED DESCRIPTION OF THE INVENTION

Described herein are peptides which may exhibit antifusogenic activity, antiviral capability, and/or

the ability to modulate intracellular processes involving coiled-coil peptide structures. peptides described include, first, DP178 (SEQ ID NO:1), a gp41-derived 36 amino acid peptide and fragments and analogs of DP178.

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In addition, the peptides of the invention described herein include peptides which are DP107 analogs. DP107 (SEQ ID NO:25) is a 38 amino acid peptide corresponding to residues 558 to 595 of the HIV-1_{LAI} transmembrane (TM) gp41 protein. Such DP107 10 analogs may exhibit antifusogenic capability, antiviral activity or an ability to modulate intracellular processes involving coiled-coil structures.

Further, peptides of the invention include DP107 and DP178 are described herein having amino acid sequences recognized by the 107x178x4, ALLMOTI5, and PLZIP search motifs. Such motifs are also discussed.

Also described here are antifusogenic, antiviral, intracellular modulatory, and diagnostic uses of the peptides of the invention. Further, procedures are 20 described for the use of the peptides of the invention for the identification of compounds exhibiting antifusogenic, antiviral or intracellular modulatory activity.

While not limited to any theory of operation, the 25 following model is proposed to explain the potent anti-HIV activity of DP178, based, in part, on the experiments described in the Examples, infra. HIV protein, gp41, DP178 corresponds to a putative α helix region located in the C-terminal end of the gp41 ectodomain, and appears to associate with a distal site on gp41 whose interactive structure is influenced

by the leucine zipper motif, a coiled-coil structure, referred to as DP107. The association of these two domains may reflect a molecular linkage or "molecular clasp" intimately involved in the fusion process. It is of interest that mutations in the C-terminal α-helix motif of gp41 (i.e., the D178 domain) tend to enhance the fusion ability of gp41, whereas mutations in the leucine zipper region (i.e., the DP107 domain) decrease or abolish the fusion ability of the viral protein. It may be that the leucine zipper motif is involved in membrane fusion while the C-terminal α-helix motif serves as a molecular safety to regulate the availability of the leucine zipper during virus-induced membrane fusion.

On the basis of the foregoing, two models are proposed of gp41-mediated membrane fusion which are 15 schematically shown in FIG. 11A-B. The reason for proposing two models is that the temporal nature of the interaction between the regions defined by DP107 and DP178 cannot, as yet, be pinpointed. Each model envisions two conformations for gp41 - one in a "native" state as it might be found on a resting The other in a "fusogenic" state to reflect virion. conformational changes triggered following binding of gp120 to CD4 and just prior to fusion with the target cell membrane. The strong binding affinity between 25 gp120 and CD4 may actually represent the trigger for the fusion process obviating the need for a pH change such as occurs for viruses that fuse within intracellular vesicles. The two major features of both models are: (1) the leucine zipper sequences (DP107) in each chain of oligomeric enveloped are held 30 apart in the native state and are only allowed access

to one another in the fusogenic state so as to form the extremely stable coiled-coils, and (2) association of the DP178 and DP107 sites as they exist in gp41 occur either in the native or fusogenic state. FIG. 11A depicts DP178/DP107 interaction in the native 5 state as a molecular clasp. On the other hand, if one assumes that the most stable form of the enveloped occurs in the fusogenic state, the model in FIG. 11B can be considered.

When synthesized as peptides, both DP107 and 10 DP178 are potent inhibitors of HIV infection and fusion, probably by virtue of their ability to form complexes with viral gp41 and interfere with its fusogenic process; e.g., during the structural transition of the viral protein from the native structure to the fusogenic state, the DP178 and DP107 15 peptides may gain access to their respective binding sites on the viral gp41, and exert a disruptive influence. DP107 peptides which demonstrate anti-HIV activity are described in Applicants' co-pending application Serial No. 08/264,531, filed June 23, 20 1994, which is incorporated by reference herein in its entirety.

As shown in the Examples, infra, a truncated recombinant gp41 protein corresponding to the ectodomain of gp41 containing both DP107 and DP178 25 domains (excluding the fusion peptide, transmembrane region and cytoplasmic domain of gp41) did not inhibit HIV-1 induced fusion. However, when a single mutation was introduced to disrupt the coiled-coil structure of the DP107 domain -- a mutation which results in a total loss of biological activity of DP107 peptides -the inactive recombinant protein was transformed to an

active inhibitor of HIV-1 induced fusion. This transformation may result from liberation of the potent DP178 domain from a molecular clasp with the leucine zipper, DP107 domain.

For clarity of discussion, the invention will be described primarily for DP178 peptide inhibitors of HIV. However, the principles may be analogously applied to other viruses, both enveloped and nonenveloped, and to other non-viral organisms.

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5.1. <u>DP178 AND DP178-LIKE PEPTIDES</u>

The DP178 peptide (SEQ ID:1) of the invention corresponds to amino acid residues 638 to 673 of the transmembrane protein gp41 from the HIV-1_{LAI} isolate, and has the 36 amino acid sequence (reading from amino to carboxy terminus):

NH2-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-COOH (SEQ ID:1)

In addition to the full-length DP178 (SEQ ID:1) 10 36-mer, the peptides of the invention may include truncations of the DP178 (SEQ ID:1) peptide which exhibit antifusogenic activity, antiviral activity and/or the ability to modulate intracellular processes involving coiled-coil peptide structures. Truncations of DP178 (SEQ ID:1) peptides may comprise peptides of 15 between 3 and 36 amino acid residues (i.e., peptides ranging in size from a tripeptide to a 36-mer polypeptide), as shown in Tables I and IA, below. Peptide sequences in these tables are listed from amino (left) to carboxy (right) terminus. "X" may represent an amino group (-NH2) and "Z" may represent a carboxyl (-COOH) group. Alternatively, "X" may represent a hydrophobic group, including but not limited to carbobenzyl, dansyl, or T-butoxycarbonyl; an acetyl group; a 9-fluorenylmethoxy-carbonyl (FMOC) 25 group; or a covalently attached macromolecular group, including but not limited to a lipid-fatty acid conjugate, polyethylene glycol, carbohydrate or peptide group. Further, "Z" may represent an amido group; a T-butoxycarbonyl group; or a covalently attached macromolecular group, including but not 30 limited to a lipid-fatty acid conjugate, polyethylene

glycol, carbohydrate or peptide group. A preferred "X" or "Z" macromolecular group is a peptide group.

TABLE I DP178 (SEO ID:1) CARBOXY TRUNCATIONS

```
5 X-YTS-Z
    X-YTSL-Z
    X-YTSLI-Z
    X-YTSLIH-Z
    X-YTSLIHS-Z
    X-YTSLIHSL-Z
    X-YTSLIHSLI-Z
    X-YTSLIHSLIE-Z
10
   X-YTSLIHSLIEE-Z
    X-YTSLIHSLIEES-Z
    X-YTSLIHSLIEESO-Z
    X-YTSLIHSLIEESON-Z
    X-YTSLIHSLIEESONO-Z
    X-YTSLIHSLIEESONOO-Z
    X-YTSLIHSLIEESQNQQE-Z
    X-YTSLIHSLIEESONOOEK-Z
15 X-YTSLIHSLIEESQNQQEKN-Z
    X-YTSLIHSLIEESQNQQEKNE-Z
    X-YTSLIHSLIEESQNQQEKNEQ-Z
    X-YTSLIHSLIEESQNOOEKNEOE-Z
    X-YTSLIHSLIEESQNOOEKNEOEL-Z
    X-YTSLIHSLIEESQNQQEKNEQELL-Z
   X-YTSLIHSLIEESQNQQEKNEOELLE-Z
    X-YTSLIHSLIEESQNQOEKNEOELLEL-Z
   X-YTSLIHSLIEESQNQQEKNEQELLELD-Z
    X-YTSLIHSLIEESQNQQEKNEOELLELDK-Z
    X-YTSLIHSLIEESQNQQEKNEQELLELDKW-Z
    X-YTSLIHSLIEESQNQQEKNEQELLELDKWA-Z
    X-YTSLIHSLIEESQNQQEKNEQELLELDKWAS-Z
    X-YTSLIHSLIEESQNQQEKNEQELLELDKWASL-Z
    X-YTSLIHSLIEESQNQQEKNEQELLELDKWASLW-Z
    X-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWN-Z
25
   X-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNW-Z
    X-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-Z
```

The one letter amino acid code is used.

TABLE IA DP178 (SEQ ID:1) AMINO TRUNCATIONS

```
X-NWF-Z
                                                    X-WNWF-Z
                                                   X-LWNWF-Z
                                                  X-SLWNWF-Z
 5
                                                 X-ASLWNWF-Z
                                                X-WASLWNWF-Z
                                               X-KWASLWNWF-Z
                                              X-DKWASLWNWF-Z
                                             X-LDKWASLWNWF-Z
                                            X-ELDKWASLWNWF-Z
                                           X-LELDKWASLWNWF-Z
                                          X-LLELDKWASLWNWF-Z
10
                                         X-ELLELDKWASLWNWF-Z
                                        X-QELLELDKWASLWNWF-Z
                                      X-EQELLELDKWASLWNWF-Z
                                     X-NEQELLELDKWASLWNWF-Z
                                    X-KNEQELLELDKWASLWNWF-Z
                                   X-EKNEQELLELDKWASLWNWF-Z
                                  X-QEKNEQELLELDKWASLWNWF-Z
                                 X-QQEKNEQELLELDKWASLWNWF-Z
15
                                X-NQQEKNEQELLELDKWASLWNWF-Z
                               X-QNQQEKNEQELLELDKWASLWNWF-Z
                              X-SQNQQEKNEQELLELDKWASLWNWF-Z
                             X-ESQNQQEKNEQELLELDKWASLWNWF-Z
                            X-EESQNQQEKNEQELLELDKWASLWNWF-Z
                           X-IEESQNQQEKNEQELLELDKWASLWNWF-Z
                          X-LIEESQNQQEKNEQELLELDKWASLWNWF-Z
                         X-SLIEESQNQQEKNEQELLELDKWASLWNWF-Z
                        X-HSLIEESQNQQEKNEQELLELDKWASLWNWF-Z
20
                       X-IHSLIEESQNQQEKNEQELLELDKWASLWNWF-Z
                      X-LIHSLIEESQNQQEKNEQELLELDKWASLWNWF-Z
                     X-SLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-Z
                    X-TSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-Z
                   X-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-Z
```

The one letter amino acid code is used.

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The peptides of the invention also include DP178like peptides. "DP178-like", as used herein, refers, first, to DP178 and DP178 truncations which contain one or more amino acid substitutions, insertions and/or deletions. Second, "DP-178-like" refers to 5 peptide sequences identified or recognized by the ALLMOTIS, 107x178x4 and PLZIP search motifs described herein, having structural and/or amino acid motif similarity to DP178. The DP178-like peptides of the invention may exhibit antifusogenic or antiviral 10 activity, or may exhibit the ability to modulate intracellular processes involving coiled-coil peptides. Further, such DP178-like peptides may possess additional advantageous features, such as, for example, increased bioavailability, and/or stability, or reduced host immune recognition.

15 HIV-1 and HIV-2 enveloped proteins are structurally distinct, but there exists a striking amino acid conservation within the DP178-corresponding regions of HIV-1 and HIV-2. The amino acid conservation is of a periodic nature, suggesting some 20 conservation of structure and/or function. Therefore, one possible class of amino acid substitutions would include those amino acid changes which are predicted to stabilize the structure of the DP178 peptides of the invention. Utilizing the DP178 and DP178 analog sequences described herein, the skilled artisan can readily compile DP178 consensus sequences and ascertain from these, conserved amino acid residues which would represent preferred amino acid substitutions.

The amino acid substitutions may be of a conserved or non-conserved nature. Conserved amino acid substitutions consist of replacing one or more

```
amino acids of the DP178 (SEQ ID:1) peptide sequence
                             amulu acids of similar charge, with amino acids of similar charge, and or
                                With amino aclos of similar charge, size, and or example, hydrophobicity characteristics, such as, for example, hydrophobicity characteristics, such as, for example, and the size of the 
                                    nydrophopicity characteristics; such as ror example, to aspartic acid (D) amino acid a glutamic acid (E) to aspartic acid (D) amino acid
                                        a gluramic acid (E) to aspartic acid (U) amino acid of consist of substitutions consist of substitution.
                                          substitution. Non-conserved substitutions consist of the DP178 (SEQ replacing one or more amino acids of the DP178 (seq
WO 01/51673
                                               replacing one or more amino acids possessing

ID:1) peptide sequence with amino acids possessing
                                                      dissimilar charge, size, and/or hydrophopically acid characteristics, such as characteristics, size, for example, a glutamic acid
                                                   dissimilar charge, size, and or hydrophobicity
                                                                               Amino acid insertions may
                                                            (E) to valine (V) substitution.
                                                                 acid residues or stretches of residues.
                                                                        Insertions may be made at the carboxy or amino peptides!
                                                                     insertions may be made at the carboxy or amino
                                                                            Lermanda emu of the peptide.

as well as at a position internal to the peptide.
                                                                               as well as at a position internal to the perture.

Such insertions will generally range from 2 to 15
                                                                                   Such in length.

The is contemplated that.
                                                                                       insertions made at either the carboxy or amino
                                                                                          terminus of the peptide of interest may be of a
                                                                                             broader size range,
                                                                                                proader size range, with about 2 to about insertions one or more such insertions acids being preferred.
                                                                                                    may be introduced into DP178 (SEQ. ID:1) or DP178
                                                                                                         truncations, as long as such insertions result in
                                                                                                               pepcides which may still be recognized by the described
                                                                                                            peptides which may still be recognized by the
                                                                                                                  NOT MAY!

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AND THE SEARCH MOULTS OF PROFIT SEARCH MOULTS O
                                                                                                                      or antivital activity
                                                                                                                         or ancivital accivity, or exhibit the ability colled-coil modulate intracellular processes involving coiled-coil
                                                                                                                                                   Preferred amino or carboxy terminal insertions
                                                                                                                                    are Pertides ranging from about 2 to about 50 amino
                                                                                                                                       are peptioes ranging from about 20 about 30 animb corresponding to gp41 protein acid residues in length,
                                                                                                                                          acla residues in length, or carboxy to the actual DP178 regions either amino to
                                                                                                                              Peptide structures.
                                                                                                                                               gp41 amino acid sequence, respectively.
```

preferred amino terminal or carboxy terminal amino acid insertion would contain gp41 amino acid sequences found immediately amino to or carboxy to the DP178 region of the gp41 protein.

Deletions of DP178 (SEQ ID:1) or DP178 truncations are also within the scope of the invention. Such deletions consist of the removal of one or more amino acids from the DP178 or DP178-like peptide sequence, with the lower limit length of the resulting peptide sequence being 4 to 6 amino acids. Such deletions may involve a single contiguous or greater than one discrete portion of the peptide sequences. One or more such deletions may be introduced into DP178 (SEQ.ID:1) or DP178 truncations, as long as such deletions result in peptides which may still be recognized by the 107x178x4, ALLMOTI5 or 15 PLZIP search motifs described herein, or may, alternatively, exhibit antifusogenic or antiviral activity, or exhibit the ability to modulate intracellular processes involving coiled-coil peptide structures.

DP178 analogs are further described, below, in Section 5.3.

5.2. DP107 AND DP107-LIKE PEPTIDES

Further, the peptides of the invention include
25 peptides having amino acid sequences corresponding to
DP107 analogs. DP107 is a 38 amino acid peptide which
exhibits potent antiviral activity, and corresponds to
residues 558 to 595 of HIV-1_{LAI} transmembrane (TM) gp41
protein, as shown here:

30 NH₂-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVERYLKDQ-COOH (SEQ ID:25)

In addition to the full-length DP107 (SEQ ID:25) 38-mer, the peptides of the invention may include truncations of the DP107 (SEQ ID:25) peptide which exhibit antifusogenic activity, antiviral activity and/or the ability to modulate intracellular processes. involving coiled-coil peptide structures. Truncations of DP107 (SEQ ID:25) peptides may comprise peptides of between 3 and 38 amino acid residues (i.e., peptides ranging in size from a tripeptide to a 38-mer polypeptide), as shown in Tables II and IIA, below. 10 Peptide sequences in these tables are listed from amino (left) to carboxy (right) terminus: "X" may represent an amino group (-NH2) and "Z" may represent a carboxyl (-COOH) group. Alternatively, "X" may represent a hydrophobic group, including but not limited to carbobenzyl, dansyl, or T-butoxycarbonyl; 15 an acetyl group; a 9-fluorenylmethoxy-carbonyl (FMOC) group; or a covalently attached macromolecular group, including but not limited to a lipid-fatty acid conjugate, polyethylene glycol, carbohydrate or peptide group. Further, "Z" may represent an amido 20 group; a T-butoxycarbonyl group; or a covalently attached macromolecular group, including but not limited to a lipid-fatty acid conjugate, polyethylene glycol, carbohydrate or peptide group. A preferred "X" or "Z" macromolecular group is a peptide group.

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TABLE II DP107 (SEO ID:25) CARBOXY TRUNCATIONS

- X-NNL-Z
- X-NNLL-Z
- X-NNLLR-Z
- 5 X-NNLLRA-Z
 - X-NNLLRAI-Z
 - X-NNLLRAIE-Z
 - X-NNLLRAIEA-Z
 - X-NNLLRAIEAQ-Z
 - X-NNLLRAIEAQQ-Z
 - X-NNLLRAIEAQOH-Z
 - X-NNLLRAIEAOOHL-Z
- 10 X-NNLLRAIEAQQHLL-Z
 - X-NNLLRAIEAOOHLLO-Z
 - X-NNLLRAIEAQOHLLOL-Z
 - X-NNLLRAIEAQQHLLQLT-Z
 - X-NNLLRAIEAOOHLLOLTV-Z
 - X-NNLLRAIEAQQHLLQLTVW-Z
 - X-NNLLRAIEAQQHLLQLTVWQ-Z
 - X-NNLLRAIEAQQHLLQLTVWOI-Z
- 15 X-NNLLRAIEAQQHLLQLTVWQIK-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQ-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQL-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQ-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQA-Z
 - X-NNLLRAIEAQQHLLQLTVWOIKOLOAR-Z
 - X-NNLLRAIEAQQHLLQLTVWOIKOLOARI-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQARIL-Z
- 20 X-NNLLRAIEAQQHLLQLTVWQIKQLQARILA-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAV-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVE-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVER-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVERY-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVERYL-Z
 - X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVERYLK-Z X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVERYLKD-Z
- 25 X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVERYLKDO-Z

The one letter amino acid code is used.

TABLE IIA DP178 (SEQ ID:25) AMINO TRUNCATIONS

	•	
	X-KDQ-	Z
	X-LKDQ-	Z
	X-YLKDQ-	Z
5	X-RYLKDQ-	Z
•	X-ERYLKDQ-	Z
	X-VERYLKDQ-	Z
	X-AVERYLKDQ-	Z
	X-LAVERYLKDQ-	Z
	X-ILAVERYLKDQ-	Z
	X-RILAVERYLKDQ-	Z
	X-ARILAVERYLKDQ-	Z
10	X-QARILAVERYLKDQ~	Z
	X-LQARILAVERYLKDQ-	Z
	X-QLQARILAVERYLKDQ-	Z
	X-KQLQARILAVERYLKDQ-	Z
	X-IKQLQARILAVERYLKDQ-	Z
		Z
	X-WQIKQLQARILAVERYLKDQ-	Z
	X-VWQIKQLQARILAVERYLKDQ-	Z
15	X-TVWQIKQLQARILAVERYLKDQ-	Z
	X-LTVWQIKQLQARILAVERYLKDQ-	\mathbf{z}
	X-QLTVWQIKQLQARILAVERYLKDQ-	Z
	X-LQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-LLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-HLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-QHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-QQHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
20	X-AQQHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-EAQQHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-IEAQQHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-AIEAQQHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-RAIEAQQHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-LRAIEAQQHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-LLRAIEAQQHLLQLTVWQIKQLQARILAVERYLKDQ-	Z
	X-NLLRAIEAQQHLLQLTVWQIKQLQARILAVERYLKDQ~	Z
25	X-NNLLRAIEAQQHLLQLTVWQIKQLQARILAVERYLKDQ-	\mathbf{z}

The one letter amino acid code is used.

The peptides of the invention also include DP107like peptides. "DP107-like", as used herein, refers, first, to DP107 and DP107 truncations which contain one or more amino acid substitutions, insertions and/or deletions. Second, "DP-107-like" refers to 5 peptide sequences identified or recognized by the ALLMOTIS, 107x178x4 and PLZIP search motifs described herein, having structural and/or amino acid motif similarity to DP107. The DP107-like peptides of the invention may exhibit antifusogenic or antiviral 10 activity, or may exhibit the ability to modulate intracellular processes involving coiled-coil peptides. Further, such DP107-like peptides may possess additional advantageous features, such as, for example, increased bioavailability, and/or stability, or reduced host immune recognition. 1.5

HIV-1 and HIV-2 enveloped proteins are structurally distinct, but there exists a striking amino acid conservation within the DP107-corresponding regions of HIV-1 and HIV-2. The amino acid conservation is of a periodic nature, suggesting some conservation of structure and/or function. Therefore, one possible class of amino acid substitutions would include those amino acid changes which are predicted to stabilize the structure of the DP107 peptides of the invention. Utilizing the DP107 and DP107 analog sequences described herein, the skilled artisan can readily compile DP107 consensus sequences and ascertain from these, conserved amino acid residues which would represent preferred amino acid substitutions.

The amino acid substitutions may be of a conserved or non-conserved nature. Conserved amino acid substitutions consist of replacing one or more

amino acids of the DP107 (SEQ ID:25) pertide sequence with amino acids of similar charge, size, and/or hydrophobicity characteristics, such as, for example, a glutamic acid (E) to aspartic acid (D) amino acid substitution. Non-conserved substitutions consist of replacing one or more amino acids of the DP107 (SEQ ID:25) pertide sequence with amino acids possessing dissimilar charge, size, and/or hydrophobicity characteristics, such as, for example, a glutamic acid (E) to valine (V) substitution.

Amino acid insertions may consist of single amino 10 acid residues or stretches of residues. The insertions may be made at the carboxy or amino terminal end of the DP107 or DP107 truncated peptides, as well as at a position internal to the peptide. Such insertions will generally range from 2 to 15 15 amino acids in length. It is contemplated that insertions made at either the carboxy or amino terminus of the peptide of interest may be of a broader size range, with about 2 to about 50 amino acids being preferred. One or more such insertions may be introduced into DP107 (SEQ.ID:25) or DP107 truncations, as long as such insertions result in peptides which may still be recognized by the 107x178x4, ALLMOTI5 or PLZIP search motifs described herein, or may, alternatively, exhibit antifusogenic 25 or antiviral activity, or exhibit the ability to modulate intracellular processes involving coiled-coil peptide structures.

Preferred amino or carboxy terminal insertions are peptides ranging from about 2 to about 50 amino acid residues in length, corresponding to gp41 protein regions either amino to or carboxy to the actual DP107 gp41 amino acid sequence, respectively. Thus, a

preferred amino terminal or carboxy terminal amino acid insertion would contain gp41 amino acid sequences found immediately amino to or carboxy to the DP107 region of the gp41 protein.

Deletions of DP107 (SEQ ID:25) or DP178 truncations are also within the scope of the invention. Such deletions consist of the removal of one or more amino acids from the DP107 or DP107-like peptide sequence, with the lower limit length of the resulting peptide sequence being 4 to 6 amino acids. 10 Such deletions may involve a single contiguous or greater than one discrete portion of the peptide sequences. One or more such deletions may be introduced into DP107 (SEO.ID:25) or DP107 truncations, as long as such deletions result in peptides which may still be recognized by the 15 107x178x4, ALLMOTI5 or PLZIP search motifs described herein, or may, alternatively, exhibit antifusogenic or antiviral activity, or exhibit the ability to modulate intracellular processes involving coiled-coil peptide structures.

DP107 and DP107 truncations are more fully described in Applicants' co-pending U.S. Patent Application Ser. No. 08/374,666, filed January 27, 1995, and which is incorporated herein by reference in its entirety. DP107 analogs are further described, below, in Section 5.3.

5.3. DP107 and DP178 ANALOGS

Peptides corresponding to analogs of the DP178, DP178 truncations, DP107 and DP107 truncation sequences of the invention, described, above, in Sections 5.1 and 5.2 may be found in other viruses,

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including, for example, non-HIV-1, enveloped viruses, non-enveloped viruses and other non-viral organisms.

The term "analog", as used herein, refers to a peptide which is recognized or identified via the 107x178x4, ALLMOTI5 and/or PLZIP search strategies 5 discussed below. Further, such peptides may exhibit antifusogenic capability, antiviral activity, or the ability to modulate intracellular processes involving coiled-coil structures.

Such DP178 and DP107 analogs may, for example, 10 correspond to peptide sequences present in TM proteins of enveloped viruses and may, additionally correspond to peptide sequences present in non enveloped and nonviral organisms. Such peptides may exhibit antifusogenic activity, antiviral activity, most particularly antiviral activity which is specific to the virus in which their native sequences are found, or may exhibit an ability to modulate intracellular processes involving coiled-coil peptide structures.

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as shown below.

DP178 analogs are peptides whose amino acid sequences are comprised of the amino acid sequences of 20 peptide regions of, for example, other (i.e., other than HIV-1_{LAI}) viruses that correspond to the gp41 peptide region from which DP178 (SEQ ID:1) was derived. Such viruses may include, but are not limited to, other HIV-1 isolates and HIV-2 isolates. 25 DP178 analogs derived from the corresponding gp41 peptide region of other (i.e., non HIV-1_{LAI}) HIV-1

NH2-YTNTIYTLLEESQNQQEKNEQELLELDKWASLWNWF-COOH 30 (DP-185; SEQ ID:3);

isolates may include, for example, peptide sequences

NH2-YTGIIYNLLEESQNQQEKNEQELLELDKWANLWNWF-COOKSEQ ID:4);

NH2-YTSLIYSLLEKSQIQQEKNEQELLELDKWASLWNWF-COOH (SEQ ID:5).

5 SEQ ID:3 (DP-185), SEQ ID:4, and SEQ ID:5 are derived from HIV-1_{SF2}, HIV-1_{RF}, and HIV-1_{MN} isolates, respectively. Underlined amino acid residues refer to those residues that differ from the corresponding position in the DP178 (SEQ ID:1) peptide. One such 10 DP178 analog, DP-185 (SEQ ID:3), is described in the Example presented in Section 6, below, where it is demonstrated that DP-185 (SEQ ID:3) exhibits antiviral activity. The DP178 analogs of the invention may also include truncations, as described above. Further, the analogs of the invention modifications such those 15 described for DP178 analogs in Section 5.1., above. It is preferred that the DP178 analogs of the invention represent peptides whose amino acid sequences correspond to the DP178 region of the gp41 protein, it is also contemplated that the peptides of the invention may, additionally, include amino sequences, ranging from about 2 to about 50 amino acid residues in length, corresponding to gp41 protein regions either amino to or carboxy to the actual DP178 amino acid sequence.

Striking similarities, as shown in FIG. 1, exist within the regions of HIV-1 and HIV-2 isolates which correspond to the DP178 sequence. A DP178 analog derived from the HIV- $2_{\scriptsize NIHZ}$ isolate has the 36 amino acid sequence (reading from amino to carboxy terminus):

³⁰ NH₂-LEANISQSLEQAQIQQEKNMYELQKLNSWDVFTNWL-COOH (SEQ ID:7)

Table III and Table IV show some possible truncations of the HIV-2_{NIHZ} DP178 analog, which may comprise peptides of between 3 and 36 amino acid residues (i.e., peptides ranging in size from a tripeptide to a 36-mer polypeptide). Peptide sequences in these 5 tables are listed from amino (left) to carboxy (right) "X" may represent an amino group (-NH,) and "Z" may represent a carboxyl (-COOH) group. Alternatively, "X" may represent a hydrophobic group, including but not limited to carbobenzyl, dansyl, or T-butoxycarbonyl; an acetyl group; a 9fluorenylmethoxy-carbonyl (FMOC) group; or a covalently attached macromolecular group, including but not limited to a lipid-fatty acid conjugate, polyethylene glycol, carbohydrate or peptide group. Further, "Z" may represent an amido group; a T-15 butoxycarbonyl group; or a covalently attached . macromolecular group, including but not limited to a lipid-fatty acid conjugate, polyethylene glycol, carbohydrate or peptide group. A preferred "X" or "Z" macromolecular group is a peptide group.

20

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TABLE III

HIV-2_{NIH2} DP178 analog carboxy truncations.

X-LEA-Z

X-LEAN-Z

X-LEANI-Z

X-LEANIS-Z

5 X-LEANISQ-Z

X-LEANISQS-Z

X-LEANISQSL-Z

X-LEANISQSLE-Z

X-LEANISQSLEQ-Z

X-LEANISQSLEQA-Z

X-LEANISQSLEQAQ-Z

X-LEANISQSLEQAQI-Z

10 X-LEANISQSLEQAQIQ-Z

X-LEANISQSLEQAQIQQ-Z

X-LEANISQSLEQAQIQQE-Z

X-LEANISQSLEQAQIQQEK-Z

X-LEANISQSLEQAQIQQEKN-Z

X-LEANISOSLEQAQIQQEKNM-Z

X-LEANISQSLEQAQIQQEKNMY-Z

X-LEANISQSLEQAQIQQEKNMYE-Z

15 X-LEANISQSLEQAQIQQEKNMYEL-Z

X-LEANISQSLEQAQIQQEKNMYELQ-Z

X-LEANISQSLEQAQIQQEKNMYELQK-Z

X-LEANISQSLEQAQIQQEKNMYELQKL-Z

X-LEANISQSLEQAQIQQEKNMYELQKLN-Z

X-LEANISQSLEQAQIQQEKNMYELQKLNS-Z

X-LEANISQSLEQAQIQQEKNMYELQKLNSW-Z

X-LEANISQSLEQAQIQQEKNMYELQKLNSWD-Z

20 X-LEANISQSLEQAQIQQEKNMYELQKLNSWDV-Z

X-LEANISQSLEQAQIQQEKNMYELQKLNSWDVF-Z

X-LEANISQSLEQAQIQQEKNMYELQKLNSWDVFT-Z

X-LEÁNISQSLEQAQIQQEKNMYELQKLNSWDVFTN-Z

X-LEANISOSLEQAQIQQEKNMYELOKLNSWDVFTNW-Z

X-LEANISQSLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z

The one letter amino acid code is used.

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X-NWL-Z X-TNWL-Z X-FTNWL-Z X-VFTNWL-Z X-DVFTNWL-Z X-WDVFTNWL-Z 5 X-SWDVFTNWL-Z X-NSWDVFTNWL-Z X-LNSWDVFTNWL-Z X-KLNSWDVFTNWL-Z X-QKLNSWDVFTNWL-Z X-LQKLNSWDVFTNWL-Z X-ELQKLNSWDVFTNWL-Z X-YELQKLNSWDVFTNWL-Z 10 X-MYELQKLNSWDVFTNWL-Z X-NMYELQKLNSWDVFTNWL-Z X-KNMYELOKLNSWDVFTNWL-Z X-EKNMYELQKLNSWDVFTNWL-Z X-QEKNMYELQKLNSWDVFTNWL-Z X-QQEKNMYELQKLNSWDVFTNWL-Z X-IQQEKNMYELQKLNSWDVFTNWL-Z X-QIQQEKNMYELQKLNSWDVFTNWL-Z 15 X-AQIQQEKNMYELQKLNSWDVFTNWL-Z ${\tt X-QAQIQQEKNMYELQKLNSWDVFTNWL-Z}$ X-EQAQIQQEKNMYELQKLNSWDVFTNWL-Z X-LEQAQIQQEKNMYELQKLNSWDVFTNWL-Z X-SLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z X-QSLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z X-SQSLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z X-ISQSLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z 20 X-NISQSLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z X-ANISQSLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z X-EANISQSLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z X-LEANISQSLEQAQIQQEKNMYELQKLNSWDVFTNWL-Z

The one letter amino acid code is used.

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DP178 and DP107 analogs are recognized or identified, for example, by utilizing one or more of the 107x178x4, ALLMOTI5 or PLZIP computer-assisted search strategies described and demonstrated, below, in the Examples presented in Sections 9 through 16 and 19 through 25. The search strategy identifies additional peptide regions which are predicted to have structural and/or amino acid sequence features similar to those of DP107 and/or DP178.

The search strategies are described fully, below, 10 in the Example presented in Section 9. While this search strategy is based, in part, on a primary amino acid motif deduced from DP107 and DP178, it is not based solely on searching for primary amino acid sequence homologies, as such protein sequence homologies exist within, but not between major groups 15 of viruses. For example, primary amino acid sequence homology is high within the TM protein of different strains of HIV-1 or within the TM protein of different isolates of simian immunodeficiency virus (SIV). Primary amino acid sequence homology between HIV-1 and 20 SIV, however, is low enough so as not to be useful. It is not possible, therefore, to find peptide regions similar to DP107 or DP178 within other viruses, or within non-viral organisms, whether structurally, or otherwise, based on primary sequence homology, alone.

Further, while it would be potentially useful to identify primary sequence arrangements of amino acids based on, for example, the physical chemical characteristics of different classes of amino acids rather than based on the specific amino acids themselves, such search strategies have, until now, proven inadequate. For example, a computer algorithm designed by Lupas et al. to identify coiled-coil

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propensities of regions within proteins (Lupas, A., et al., 1991 Science <u>252</u>:1162-1164) is inadequate for identifying protein regions analogous to DP107 or DP178.

Specifically, analysis of HIV-1 gp160 (containing both gp120 and gp41) using the Lupas algorithm does not identify the coiled-coil region within Dp107. It does, however, identify a region within Dp178 beginning eight amino acids N-terminal to the start of Dp178 and ending eight amino acids from the C-terminus. The Dp107 peptide has been shown experimentally to form a stable coiled coil. A search

based on the Lupas search algorithm, therefore, would not have identified the DP107 coiled-coil region.

Conversely, the Lupas algorithm identified the DP178 region as a potential coiled-coil motif. However, the peptide derived from the DP178 region failed to form a coiled coil in solution.

A possible explanation for the inability of the Lupas search algorithm to accurately identify coiled-coil sequences within the HIV-1 TM, is that the Lupas algorithm is based on the structure of coiled coils from proteins that are not structurally or functionally similar to the TM proteins of viruses, antiviral peptides (e.g. DP107 and DP178) of which are an object of this invention.

The computer search strategy of the invention, as demonstrated in the Examples presented below, in Sections 9 through 16 and 19 through 25, successfully identifies regions of proteins similar to DP107 or DP178. This search strategy was designed to be used with a commercially-available sequence database package, preferably PC/Gene.

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A series of search motifs, the 107x178x4, ALLMOTIS and PLZIP motifs, were designed and engineered to range in stringency from strict to broad, as discussed in this Section and in Section 9, with 107x178x4 being preferred. The sequences 5 identified via such search motifs, such as those listed in Tables V-XIV, below, potentially exhibit antifusogenic, such as antiviral, activity, may additionally be useful in the identification of antifusogenic, such as antiviral, compounds, and are intended to be within the scope of the invention.

Coiled-coiled sequences are thought to consist of heptad amino acid repeats. For ease of description, the amino acid positions within the heptad repeats are sometimes referred to as A through G, with the first position being A, the second B, etc. The motifs used to identify DP107-like and DP178-like sequences herein are designed to specifically search for and identify such heptad repeats. In the descriptions of each of the motifs described, below, amino acids enclosed by brackets , i.e., [], designate the only amino acid residues that are acceptable at the given position, while amino acids enclosed by braces, i.e., {}, designate the only amino acids which are unacceptable at the given heptad position. When a set of bracketed or braced amino acids is followed by a number in 25 parentheses i.e., (), it refers to the number of subsequent amino acid positions for which the designated set of amino acids hold, e.g, a (2) means "for the next two heptad amino acid positions".

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The ALLMOTI5 is written as follows:

```
{CDGHP}-{CFP}(2)-{CDGHP}-{CFP}(3)-
{CDGHP}-{CFP}(2)-{CDGHP}-{CFP}(3)-
{CDGHP}-{CFP}(2)-{CDGHP}-{CFP}(3)-
{CDGHP}-{CFP}(2)-{CDGHP}-{CFP}(3)-
{CDGHP}-{CFP}(2)-{CDGHP}-{CFP}(3)-
```

Translating this motif, it would read: "at the 5 first (A) position of the heptad, any amino acid residue except C, D, G, H, or P is acceptable, at the next two (B,C) amino acid positions, any amino acid residue except C, F, or P is acceptable, at the fourth heptad position (D), any amino acid residue except C, D, G, H, or P is acceptable, at the next three (E, F, G) amino acid positions, any amino acid residue except C, F, or P is acceptable. This motif is designed to search for five consecutive heptad repeats (thus the repeat of the first line five times), meaning that it searches for 35-mer sized peptides. It may also be designed to search for 28-mers, by only repeating the initial motif four times. With respect to the ALLMOTI5 motif, a 35-mer search is preferred. viral (non-bacteriophage) sequences identified via such an ALLMOTI5 motif are listed in Table V in U.S. 20 Patent Application No. 08/470,896 filed on June 6, 1995 which is incorporated herein by reference in its entirety. These viral sequences potentially exhibit antiviral activity, may be useful in the the identification of antiviral compounds, and are 25 intended to be within the scope of the invention. those instances wherein a single gene exhibits greater than one sequence recognized by the ALLMOTIS search motif, the amino a cid residue numbers of these sequences are listed under "Area 2", Area 3", etc. This convention is used for each of the Tables listed, below, at the end of this Section.

The 107x178x4 motif is written as follows:

```
[EFIKLNQSTVWY] - {CFMP} (2) - [EFIKLNQSTVWY] - {CFMP} (3) -
[EFIKLNQSTVWY] - {CFMP} (2) - [EFIKLNQSTVWY] - {CFMP} (3) -
[EFIKLNQSTVWY] - {CFMP} (2) - [EFIKLNQSTVWY] - {CFMP} (3) -
[EFIKLNQSTVWY] - {CFMP} (2) - [EFIKLNQSTVWY] - {CFMP} (3) -
```

Translating this motif, it would read: "at the first (A) position of the heptad, only amino acid residue E, F, I, K, L, N, Q, S, T, V, W, or Y is acceptable, at the next two (B,C) amino acid positions, any amino acid residue except C, F, M or P is acceptable, at the fourth position (D), only amino acid residue E, F, I, K, L, N, Q, S, T, V, W, or Y is acceptable, at the next three (E, F, G) amino acid positions, any amino acid residue except C, F, M or P is acceptable. This motif is designed to search for four consecutive heptad repeats (thus the repeat of the first line four times), meaning that it searches 15 for 28-mer sized peptides. It may also be designed to search for 35-mers, by repeating the initial motif five times. With respect to the 107x178x4 motif, a 28-mer search is preferred.

Those viral (non-bacteriophage) sequences

identified via such a 107x178x4 motif are listed in
Table VI in U.S. Patent Application No. 08/470,896
filed on June 6, 1995, which is incorporated herein,
by reference, in its entirety. Those viral (nonbacteriophage) sequences listed in Table VII of U.S.

Patent Application No. 08/470,896 (incorporated herein
by reference in its entirety) are particularly
preferred.

The 107x178x4 search motif was also utilized to identify non-viral procaryotic protein sequences, as listed in Table VIII in U.S. Patent Application No. 08/470,896 filed on June 6, 1995, which is incorporated herein, by reference, in its entirety.

Further, this search motif was used to reveal a number of human proteins. The results of this human protein 107x178x4 search is listed in Table IX in U.S. Patent Application No. 08/470,896 filed on June 6, 1995, which is incorporated herein, by reference, in its entirety. The sequences listed in Tables VIII and IX, therefore, reveal peptides which may be useful as antifusogenic compounds or in the identification of antifusogenic compounds, and are intended to be within the scope of the invention.

The PLZIP series of motifs are as listed in FIG. 10 These motifs are designed to identify leucine zipper coiled-coil like heptads wherein at least one proline residue is present at some predefined distance N-terminal to the repeat. These PLZIP motifs find regions of proteins with similarities to HIV-1 DP178 15 generally located just N-terminal to the transmembrane anchor. These motifs may be translated according to the same convention described above. Each line depicted in FIG. 19 represents a single, complete search motif. "X" in these motifs refers to any amino acid residue. In instances wherein a motif contains two numbers within parentheses, this refers to a variable number of amino acid residues. For example, X (1,12) is translated to "the next one to twelve amino acid residues, inclusive, may be any amino 25 acid".

Tables X through XIV in U.S. Patent Application No. 08/470,896 filed on June 6, 1995 (which is incorporated herein, by reference, in its entirety), list sequences identified via searches conducted with such PLZIP motifs. Specifically, Table X lists viral sequences identified via PCTLZIP, P1CTLZIP and P2CTLZIP search motifs, Table XI lists viral sequences

identified via P3CTLZIP, P4CTLZIP, P5CTLZIP and
P6CTLZIP search motifs, Table XII lsts viral sequences
identified via P7CTLZIP, P8CTLZIP and P9CTLZIP search
motifs, Table XIII lists viral sequences identified
via P12LZIPC searches and Table XIV lists viral

sequences identified via P23TLZIPC search motifs The
viral sequences listed in these tables represent
peptides which potentially exhibit antiviral activity,
may be useful in the identification of antiviral
compounds, and are intended to be within the scope of
the invention.

The Examples presented in Sections 17, 18, 26 and 27 below, demonstrate that viral sequences identified via the motif searches described herein identify substantial antiviral characteristics. Specifically, the Example presented in Section 17 describes peptides with anti-respiratory syncytial virus activity, the Example presented in Section 18 describes peptides with anti-parainfluenza virus activity, the Example presented in Section 26 describes peptides with anti-measles virus activity and the Example presented in Section 27 describes peptides with anti-simian immunodeficiency virus activity.

The DP107 and DP178 analogs may, further, contain any of the additional groups described for DP178, above, in Section 5.1. For example, these peptides

25 may include any of the additional amino-terminal groups as described above for "X" groups, and may also include any of the carboxy-terminal groups as described, above, for "Z" groups.

Additionally, truncations of the identified DP107 and DP178 peptides are among the peptides of the invention. Further, such DP107 and DP178 analogs and DP107/DP178 analog truncations may exhibit one or more

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amino acid substitutions, insertion, and/or deletions.
                                                                                                                         amino acia substitutions; insertions insertions

The DP178 analog amino acid substitutions;
                                                                                                                                          The DELIGIB analog amino acid substitutions, ppi78-like for ppi78-like above, for ppi78-like and deletions, are as described, and deletions, 
                                                                                                                                                      and delections, are as described, above, for DP178-like

and delections, are as described, above, for DP178-like

The DP-107 analog amino acid

peptides in Section 5.1.
                                                                                                                                                                    pepciaes in section 3.1. The uk-lul analog amino aci
                                                                                                                                                                                substitutions, insertions and deletions are also as in Section described, above, above
WO 01/51673
                                                                                                                                                                                                 The perfective examples of such DP107/DP178

5.2.

Representative examples of such DP107/DP178
                                                                                                                                                                                                             truncations are provided in Tables W through XXII of truncations are provided on the part of the part 
                                                                                                                                                                                                                         truncations are provided in Tables XV through XXII of through XXII of the through XXII of through XXII of the through XXII of 
                                                                                                                                                                                                                                            June 6: 1995; which is incorporated herein by
                                                                                                                                                                                                                                                                                                                                       Other exemplary DP178 and DP107 Peptides and
                                                                                                                                                                                                                                                                                            DP178-like and DP107-like peptides wnlch are include the present invention include serial considered part of the present annication serial considered part in a parent annication
                                                                                                                                                                                                                                                                                DP178-like and DP107-like Peptides which are
                                                                                                                                                                                                                                                           reference in its entirety.
                                                                                                                                                                                                                                                                                                          considered part of the present Application serial

U.S. Patent Application

peptides described in U.S. Patent Application
                                                                                                                                                                                                                                                                                                                         No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315,304 filed on May 4, 1999 which is No. 09/315, 1999 
                                                                                                                                                                                                                                                                                                                                     incorporated by reference in its entirety.
                                                                                                                                                                                                                                                                                                                                                        and DP107 peptides and DP178-like
                                                                                                                                                                                                                                                                                                                                                                 and writes include.

peptides include.

peptides include.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Other DP178, DP107, DP178-like and DP107-like
                                                                                                                                                                                                                                                                                                                                                   Other DP178, DP107, DP178-like and Ukiui-like

Other Dp178, DP107, DP178-like and Ukiui-like

10.5.
                                                                                                                                                                                                                                                                                                                                                                                                                             perunes muchane perunes aescribed e.g., filed on on on one and application serial No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                     Warch 29, 1993, now U.S. serial No. 08/073,028 filed on U.S. serial No. 08/073,028 in To
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Object, 101 riled on Serial No. 08/470, 896 filed on June
Patent Application Serial information in incommentation in incomments 
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         racent Application serial Mo. volting herein by
6, 1995 each of which is incorporated herein by
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         reference in its entirety.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          _ 52 -
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TABLE V

		TABLE V
	T	Carriage
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	9	RQLLSGIVQQQNNLLRAIEAQQHLLQLT
10 15 20	10	MTLTVQARQLLSGIVQQQNNLLRAIEAQ
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          Ac-ISTELGNVNNSISNALDKLEESNSKLDKVNVKLTS-NH2
    277
    278
          Ac-DISTELGNVNNSISNALDKLEESNSKLDKVNVKLT-NH2
    279
          Ac-LDISTELGNVNNSISNALDKLEESNSKLDKVNVKL-NH2
    280
          Ac-NLDISTELGNVNNSISNALDKLEESNSKLDKVNVK-NH2
    281
          Ac-GNLDISTELGNVNNSISNALDKLEESNSKLDKVNV-NH2
    282
          Ac-TGNLDISTELGNVNNSISNALDKLEESNSKLDKVN-NH2
    283
          Ac-VTGNLDISTELGNVNNSISNALDKLEESNSKLDKV-NH2
    284
          Ac-IVTGNLDISTELGNVNNSISNALDKLEESNSKLDK-NH2
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          AC-VIVTGNLDISTELGNVNNSISNALDKLEESNSKLD-NH2
    286
          Ac-QVIVTGNLDISTELGNVNNSISNALDKLEESNSKL-NH2
    287
          Ac-SQVIVTGNLDISTELGNVNNSISNALDKLEESNSK-NH2
    288
          Ac-DSQVIVTGNLDISTELGNVNNSISNALDKLEESNS-NH2
    289
          Ac-LDSQVIVTGNLDISTELGNVNNSISNALDKLEESN-NH2
    290
          Ac-ILDSQVIVTGNLDISTELGNVNNSISNALDKLEES-NH2
    291
          Ac-SILDSQVIVTGNLDISTELGNVNNSISNALDKLEE-NH2
          Ac-ISILDSQVIVTGNLDISTELGNVNNSISNALDKLE-NH2
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    293
          Ac-NISILDSQVIVTGNLDISTELGNVNNSISNALDKL-NH2
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    294
         . Ac-KNISILDSQVIVTGNLDISTELGNVNNSISNALDK-NH2
    295
          Ac-QKNISILDSQVIVTGNLDISTELGNVNNSISNALD-NH2
          Ac-YQKNISILDSQVIVTGNLDISTELGNVNNSISNAL-NH2
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    297
          Ac-TYQKNISILDSQVIVTGNLDISTELGNVNNSISNA-NH2
    298
          Ac-ATYQKNISILDSQVIVTGNLDISTELGNVNNSISN-NH2
    299
          Ac-DATYQKNISILDSQVIVTGNLDISTELGNVNNSIS-NH2
          Ac-FDATYQKNISILDSQVIVTGNLDISTELGNVNNSI-NH2
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          Ac-EFDATYQKNISILDSQVIVTGNLDISTELGNVNNS-NH2
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    302
          Ac-GEFDATYQKNISILDSQVIVTGNLDISTELGNVNN-NH2
    303
          Ac-SGEFDATYQKNISILDSQVIVTGNLDISTELGNVN-NH2
    304
          Ac-LSGEFDATYQKNISILDSQVIVTGNLDISTELGNV-NH2
          Ac-RLSGEFDATYQKNISILDSQVIVTGNLDISTELGN-NH2
    305
          Ac-LRLSGEFDATYQKNISILDSQVIVTGNLDISTELG-NH2
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    307
          Ac-TLRLSGEFDATYQKNISILDSQVIVTGNLDISTEL-NH2
    308
          AC-ITLRLSGEFDATYQKNISILDSQVIVTGNLDISTE-NH2
          Ac-GITLRLSGEFDATYQKNISILDSQVIVTGNLDIST-NH2
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          Ac-TATIEAVHEVTDGLSQLAVAVGKMQQFVNDQFNNT-NH2
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    311
          AC-ITATIEAVHEVTDGLSQLAVAVGKMQOFVNDOFNN-NH2
    312
          AC-SITATIEAVHEVTDGLSQLAVAVGKMQOFVNDOFN-NH2
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No.
          Sequence
    314
          Ac-KESITATIEAVHEVTDGLSQLAVAVGKMQQFVNDQ-NH2
          Ac-LKESITATIEAVHEVTDGLSOLAVAVGKMQOFVND-NH2
    315
          Ac-RLKESITATIEAVHEVTDGLSQLAVAVGKMQQFVN-NH2
    316
    317
          Ac-LRLKESITATIEAVHEVTDGLSQLAVAVGKMQQFV-NH2
    318
          Ac-ILRLKESITATIEAVHEVTDGLSQLAVAVGKMQQF-NH2
    319
          Ac-NILRLKESITATIEAVHEVTDGLSQLAVAVGKMQQ-NH2
          Ac-ANILRLKESITATIEAVHEVTDGLSQLAVAVGKMQ-NH2
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    321
          Ac-AANILRLKESITATIEAVHEVTDGLSQLAVAVGKM-NH2
    322
          Ac-HKCDDECMNSVKNGTYDYPKYEEESKLNRNEIKGV-NH2
          Ac-KCDDECMNSVKNGTYDYPKYEEESKLNRNEIKGVK-NH2
    324
          Ac-CDDECMNSVKNGTYDYPKYEEESKLNRNEIKGVKL-NH2
          Ac-DDECMNSVKNGTYDYPKYEEESKLNRNEIKGVKLS-NH2
    325
    326
          Ac-DECMNSVKNGTYDYPKYEEESKLNRNEIKGVKLSS-NH2
    327
          Ac-ECMNSVKNGTYDYPKYEEESKLNRNEIKGVKLSSM-NH2
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    328
          Ac-CMNSVKNGTYDYPKYEEESKLNRNEIKGVKLSSMG-NH2
    329
          Ac-MNSVKNGTYDYPKYEEESKLNRNEIKGVKLSSMGV-NH2
          Ac-NSVKNGTYDYPKYEEESKLNRNEIKGVKLSSMGVY-NH2
    330
          Ac-SVKNGTYDYPKYEEESKLNRNEIKGVKLSSMGVYQ-NH2
    331
    332
          Ac-VKNGTYDYPKYEEESKLNRNEIKGVKLSSMGVYQI-NH2
          Ac-KNGTYDYPKYEEESKLNRNEIKGVKLSSMGVYQIL-NH2
    334
          Ac-AFIRKSDELLHNV-NH2
    335
          Ac-VVLAGAALGVATAAQITAGIALHOSMLNSQAIDNL-NH2
          Ac-VLAGAALGVATAAQITAGIALHQSMLNSQAIDNLR-NH2
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    337
          Ac-LAGAALGVATAAQITAGIALHQSMLNSQAIDNLRA-NH2
    338
          Ac-AGAALGVATAAQITAGIALHOSMLNSQAIDNLRAS-NH2
    339
          Ac-GAALGVATAAQITAGIALHQSMLNSQAIDNLRASL-NH2
    340
          Ac-AALGVATAAOITAGIALHOSMLNSOAIDNLRASLE-NH2
    341
          Ac-ALGVATAAQITAGIALHQSMLNSQAIDNLRASLET-NH2
    342
          Ac-LGVATAAQITAGIALHQSMLNSQAIDNLRASLETT-NH2
    343
          Ac-GVATAAQITAGIALHQSMLNSQAIDNLRASLETTN-NH2
          Ac-VATAAQITAGIALHQSMLNSQAIDNLRASLETTNQ-NH2
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    345
          Ac-ATAAQITAGIALHQSMLNSQAIDNLRASLETTNQA-NH2
    346
          Ac-TAAOITAGIALHOSMLNSOAIDNLRASLETTNOAI-NH2
          Ac-AAQITAGIALHQSMLNSQAIDNLRASLETTNQAIE-NH2
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          Ac-AQITAGIALHQSMLNSQAIDNLRASLETTNQAIEA-NH2
    349
          Ac-QITAGIALHQSMLNSQAIDNLRASLETTNQAIEAI-NH2
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          Ac-ITAGIALHQSMLNSQAIDNLRASLETTNQAIEAIR-NH2
    351
          Ac-TAGIALHQSMLNSQAIDNLRASLETTNQAIEAIRQ-NH2
    352
          Ac-AGIALHOSMLNSOAIDNLRASLETTNOAIEAIROA-NH2
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    353
          Ac-GIALHQSMLNSQAIDNLRASLETTNQAIEAIRQAG-NH2
    354
          Ac-IALHQSMLNSQAIDNLRASLETTNQAIEAIRQAGQ-NH2
    355
          Ac-ALHQSMLNSQAIDNLRASLETTNQAIEAIRQAGQE-NH2
    356
           Ac-LHQSMLNSQAIDNLRASLETTNQAIEAIRQAGQEM-NH2
     357
          Ac-HQSMLNSQAIDNLRASLETTNOAIEAIROAGOEMI-NH2
     358
          Ac-QSMLNSQAIDNLRASLETTNQAIEAIRQAGQEMIL-NH2
     359
           Ac-SMLNSQAIDNLRASLETTNQAIEAIRQAGQEMILA-NH2
     360
          Ac-MLNSQAIDNLRASLETTNQAIEAIRQAGQEMILAV-NH2
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     361
          Ac-LNSQAIDNLRASLETTNQAIEAIRQAGQEMILAVQ-NH2
     362
           Ac-NSQAIDNLRASLETTNQAIEAIRQAGQEMILAVQG-NH2
          Ac-SQAIDNLRASLETTNOAIEAIROAGOEMILAVOGV-NH2
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          Sequence
    No.
    364
          Ac-QAIDNLRASLETTNQAIEAIRQAGQEMILAVQGVQ-NH2
          Ac-AIDNLRASLETTNQAIEAIRQAGQEMILAVQGVQD-NH2
    366
          Ac - IDNLRASLETTNQAIEAIRQAGQEMILAVQGVQDY-NH2
    367
          Ac-DNLRASLETTNOAIEAIROAGOEMILAVOGVODYI-NH2
          AC-NLRASLETTNQAIEAIRQAGOEMILAVOGVODYIN-NH2
    368
          Ac-LRASLETTNQAIEAIRQAGQEMILAVQGVQDYINN-NH2
    369
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    370
          Ac-RASLETTNQAIEAIRQAGQEMILAVQGVQDYINNE-NH2
          AC-YTSVITIELSNIKENKUNGTDAVKLIKOELDKYK-NH2
    371
          Ac-TSVITIELSNIKENKUNGTDAVKLIKQELDKYKN-NH2
    372
          Ac-SVITIELSNIKENKUNGTDAVKLIKOELDKYKNA-NH2
    373
          Ac-SNIKENKUNGTDAKVKLIKOELDKYKNAVTELOLL-NH2
    374
          Ac-KENKUNGTDAKVKLIKQELDKYKNAVTELQLLMQS-NH2
    375
          Ac-CLELDKWASLWNWFC-NH2
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    377
          Ac-CLELDKWASLANWFC-NH2
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    378
          Ac-CLELDKWASLFNFFC-NH2
          Ac-YTSLIHSLIEESQNQQEKNEQELLELDKWASLFNFF-NH2
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    381
          Ac-RMKQLEDKVEELLSKNYHLENELELDKWASLWNWF-NH2
    382
          AC-KVEELLSKNYHLENELELDKWASLWNWF-NH2
          Ac-RMKQLEDKVEELLSKLEWIRRSNQKLDSI-NH2
    384
          AC-RMKOLEDKVEELLSKLAFIRKSDELLHNV-NH2
    385
          Ac-ELEALRGELRALRGELELDKWASLWNWF-NH2
    386
          Ac-LDPIDISIELNKAKSDLEESKEWIRRSNQKLDSI-NH2
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    387
          Ac-CNEQLSDSFPVEFFQV-NH2
          Ac-MAEDDPYLGRPEQMFHLDPSL-NH2
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    389
          Ac-EDFSSIADMDFSALLSOISS-NH2
          Ac-TWQEWERKVDFLEENITALLEEAQIQQEKNMYELQ-NH2
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    391
          Ac-WQEWERKVDFLEENITALLEEAQIQQEKNMYELQK-NH2
    392
          Ac-QEWERKVDFLEENITALLEEAQIQQEKNMYELQKL-NH2
    393
          Ac-EWERKVDFLEENITALLEEAQIQQEKNMYELQKLN-NH2
    394
          Ac-WERKVDFLEENITALLEEAQIQQEKNMYELQKLNS-NH2
    395
          Ac-ERKVDFLEENITALLEEAOIOOEKNMYELOKLNSW-NH2
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          Ac-RKVDFLEENITALLEEAQIQOEKNMYELOKLNSWD-NH2
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    397
          Ac-KVDFLEENITALLEEAQIQQEKNMYELQKLNSWDV-NH2
    398
          Ac-VDFLEENITALLEEAQIQQEKNMYELQKLNSWDVF-NH2
    399
          Ac-DFLEENITALLEEAQIQQEKNMYELQKLNSWDVFG-NH2
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    402
          Ac-LEENITALLEEAQIQQEKNMYELQKLNSWDVFGNWF-NH2
    403
          Ac-NEQSEEKENELYWAKEQLLDLLFNIFNQTVGAWIMQ-NH2
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    406
           Ac-QQLLDVVKRQQELLRLTVWGTKNLQTRVTAIEKYLKDQ-NH2
    407
          Ac-QQLLDVVKRQQELLRLTVWGPKNLQTRVTAIEKYLKDQ-NH2
    408
          Ac-DERKQDKVLVVQQTGTLQLTLIQLEKTAKLQWVRLNRY-NH2
           Ac-QQQLLDVVKRQQELLRLTVWGTKNLQTRVTAIEKY-NH2
    409
           Ac-QQLLDVVKRQQELLRLTVWGTKNLQTRVTAIEKYL-NH2
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    411
           Ac-QLLDVVKRQQELLRLTVWGTKNLQTRVTAIEKYLK-NH2
     412
           Ac-LLDVVKRQQELLRLTVWGTKNLQTRVTAIEKYLKD-NH2
    413
           Ac-LDVVKRQQELLRLTVWGTKNLOTRVTAIEKYLKDO-NH2
           Ac-DVVKRQQELLRLTVWGTKNLQTRVTAIEKYLKDQA-NH2
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           Ac-VVKRQQELLRLTVWGTKNLQTRVTAIEKYLKDQAQ-NH2
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No.
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          Ac-VKRQQELLRLTVWGTKNLQTRVTAIEKYLKDQAQL-NH2
    416
          Ac-KRQQELLRLTVWGTKNLQTRVTAIEKYLKDQAQLN-NH2
    417
          Ac-ROOELLRLTVWGTKNLQTRVTAIEKYLKDQAQLNA-NH2
    418
          Ac-QQELLRLTVWGTKNLQTRVTAIEKYLKDQAQLNAW-NH2
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    420
          Ac-QELLRLTVWGTKNLQTRVTAIEKYLKDQAQLNAWG-NH2
          Ac-ELLRLTVWGTKNLQTRVTAIEKYLKDQAQLNAWGC-NH2
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    422
          Ac-NNLLRAIEAOOHLLQLTVWGPKQLQARILAVERYLKDQ-NH2
          Ac-SELEIKRYKNRVASRKCRAKFKQLLQHYREVAAAK-NH2
    423
    424
          Ac-ELEIKRYKNRVASRKCRAKFKQLLQHYREVAAAKS-NH2
    425
          Ac-LEIKRYKNRVASRKCRAKFKQLLQHYREVAAAKSS-NH2
          Ac-EIKRYKNRVASRKCRAKFKQLLQHYREVAAAKSSE-NH2
    426
          Ac-IKRYKNRVASRKCRAKFKQLLQHYREVAAAKSSEN-NH2
    427
    428
          Ac-KRYKNRVASRKCRAKFKQLLQHYREVAAAKSSEND-NH2
          Ac-RYKNRVASRKCRAKFKQLLQHYREVAAAKSSENDR-NH2
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     430
          Ac-YKNRVASRKCRAKFKOLLOHYREVAAAKSSENDRL-NH2
    431
          Ac-KNRVASRKCRAKFKQLLQHYREVAAAKSSENDRLR-NH2
    432
          Ac-NRVASRKCRAKFKQLLQHYREVAAAKSSENDRLRL-NH2
          Ac-RVASRKCRAKFKQLLQHYREVAAAKSSENDRLRLL-NH2
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     434
          Ac-VASRKCRAKFKOLLOHYREVAAAKSSENDRLRLLL-NH2
     435
          Ac-ASRKCRAKFKQLLQHYREVAAAKSSENDRLRLLLK-NH2
     436
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     437
           Ac-RKCRAKFKQLLQHYREVAAAKSSENDRLRLLLKQM-NH2
    438
           Ac-KCRAKFKQLLQHYREVAAAKSSENDRLRLLLKOMC-NH2
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           Ac-CRAKFKQLLQHYREVAAAKSSENDRLRLLLKOMCP-NH2
    440
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     441
           Ac-AKFKQLLQHYREVAAAKSSENDRLRLLLKQMCPSL-NH2
           Ac-KFKQLLQHYREVAAAKSSENDRLRLLLKQMCPSLD-NH2
     443
           Ac-FKOLLOHYREVAAAKSSENDRLRLLLKOMCPSLDV-NH2
     444
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     445
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     446
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           Ac-LQHYREVAAAKSSENDRLRLLLKQMCPSLDVDSII-NH2
     448
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     449
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     451
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           Ac-EVAAAKSSENDRLRLLLKQMCPSLDVDSIIPRTPD-NH2
     453
           Ac-VAAAKSSENDRLRLLLKQMCPSLDVDSIIPRTPDV-NH2
     454
           Ac-AAAKSSENDRLRLLLKQMCPSLDVDSIIPRTPDVL-NH2
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           Ac-AAKSSENDRLRLLLKQMCPSLDVDSIIPRTPDVLH-NH2
     456
           Ac-AKSSENDRLRLLLKQMCPSLDVDSIIPRTPDVLHE-NH2
     457
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           Ac-SSENDRLRLLLKQMCPSLDVDSIIPRTPDVLHEDL-NH2
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     459
           Ac-SENDRLRLLLKQMCPSLDVDSIIPRTPDVLHEDLL-NH2
     460
           Ac-ENDRLRLLLKQMCPSLDVDSIIPRTPDVLHEDLLN-NH2
     461
           Ac-NDRLRLLLKQMCPSLDVDSIIPRTPDVLHEDLLNF-NH2
           Ac-PGYRWMCLRRFIIFLFILLLCLIFLLVLLDYOGML-NH2
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           Ac-GYRWMCLRRFIIFLFILLLCLIFLLVLLDYQGMLP-NH2
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     536
           Ac-YRWMCLRRFIIFLFILLCLIFLLVLLDYQGMLPV-NH2
     537
           Ac-RWMCLRRFIIFLFILLLCLIFLLVLLDYOGMLPVC-NH2
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No.
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    538
           Ac-WMCLRRFIIFLFILLLCLIFLLVLLDYQGMLPVCP-NH2
    539
           Ac-MCLRRFIIFLFILLLCLIFLLVLLDYOGMLPVCPL-NH2
           Ac-CLRRFIIFLFILLLCLIFLLVLLDYOGMLPVCPLI-NH2
    541
           Ac-LRRFIIFLFILLLCLIFLLVLLDYOGMLPVCPLIP-NH2
          Ac-RRFIIFLFILLCLIFLLVLLDYQGMLPVCPLIPG-NH2
    542
     543
           Ac-RFIIFLFILLLCLIFLLVLLDYQGMLPVCPLIPGS-NH2
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    544
           Ac-FIIFLFILLCLIFLLVLLDYQGMLPVCPLIPGSS-NH2
           Ac-IIFLFILLCLIFLLVLLDYQGMLPVCPLIPGSST-NH2
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    546
          Ac-IFLFILLCLIFLLVLLDYQGMLPVCPLIPGSSTT-NH2
    547
          Ac-FLFILLCLIFLLVLLDYQGMLPVCPLIPGSSTTS-NH2
    548
          Ac-LFILLCLIFLLVLLDYQGMLPVCPLIPGSSTTST-NH2
           Ac-FILLLCLIFLLVLLDYQGMLPVCPLIPGSSTTSTG-NH2
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          Ac-ILLLCLIFLLVLLDYQGMLPVCPLIPGSSTTSTGP-NH2
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           Ac-LLLCLIFLLVLLDYQGMLPVCPLIPGSSTTSTGPC-NH2
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    552
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     553
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    554
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    555
           Ac-LIFLLVLLDYQGMLPVCPLIPGSSTTSTGPCRTCM-NH2
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    557
           Ac-FLLVLLDYQGMLPVCPLIPGSSTTSTGPCRTCMTT-NH2
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    559
          Ac-LLVLQAGFFLLTRILTIPQSLDSWWTSLNFLGGTT-NH2
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           Ac-LVLQAGFFLLTRILTIPQSLDSWWTSLNFLGGTTV-NH2
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    562
          Ac-LQAGFFLLTRILTIPQSLDSWWTSLNFLGGTTVCL-NH2
    563
          Ac-QAGFFLLTRILTIPQSLDSWWTSLNFLGGTTVCLG-NH2
    564
          Ac-AGFFLLTRILTIPQSLDSWWTSLNFLGGTTVCLGQ-NH2
    565
          Ac-GFFLLTRILTIPQSLDSWWTSLNFLGGTTVCLGON-NH2
    566
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          Ac-FLLTRILTIPQSLDSWWTSLNFLGGTTVCLGQNSQ-NH2
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    568
          Ac-LLTRILTIPQSLDSWWTSLNFLGGTTVCLGQNSOS-NH2
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    569
          Ac-LTRILTIPQSLDSWWTSLNFLGGTTVCLGQNSQSP-NH2
    570
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    571
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          Ac-CGGNNLLRAIEAQQHLLQLTVWGIKQLQARILAVERYLKDQ-NH2
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    575
          AC-AVSKGYLSALRTGWYTSVITIELSNIKENKUNGTDA-NH2
          Ac-SISNIETVIEFQQKNNRLLEITREFSVNAGVTTPVS-NH2
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    577
          Ac-DQQIKQYKRLLDRLIIPLYDGLRQKDVIVSNQESN-NH2
    578
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    579
          Ac-TSITLQVRLPLLTRLLNTQIYRVDSISYNIONREWY-NH2
          Ac-VEIAEYRRLLRTVLEPIRDALNAMTQNIRPVQSVA-NH2
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          Ac-LKEAIRDTNKAVQSVQSSIGNLIVAIKS-NH2
    583
          NNLLRAIEAQQHLLQLTVWGIKQLQARILAVERYLKDO-NH2
    583
          NNLLRAIEAQQHLLQLTVWGIKQLQARILAVERYLKDQ-NH2
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    584
           QKQEPIDKELYPLTSL
    585
          YPKFVKONTLKLAT
    586
          QYIKANQKFIGITE
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    No.
           Sequence
    587
          NGOIGNDPNRDILY
    588
          AC-RPDVY-OH
          CLELDKWASLWNWFC-(cyclic)
    589
          CLELDKWASLANWFC-(cyclic)
    590
    591
           CLELDKWASLANFFC-(cyclic)
    594
          Ac-NNLLRAIEAQQQHLLQLTVWGIKQLQARILAVERYLKDQ-NH2
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          Ac-CGGYTSLIHSLIEESONOOEKNEOELLELDKWASLWNNWF-NH2
     596
          Ac-PLLVLOAGFFLLTRILTIPOSLDSWWTSLNFLGGT-NH2
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          Ac-LLVLQAGFFLLTRILTIPQSLDSWWTSLNFLGGTT-NH2
    598
          Ac-LVLQAGFFLLTRILTIPQSLDSWWTSLNFLGGTTV-NH2
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           Ac-VLQAGFFLLTRILTIPQSLDSWWTSLNFLGGTTVC-NH2
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          Ac-LOAGFFLLTRILTIPOSLDSWWTSLNFLGGTTVCL-NH2
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           Ac-LTRILTIPQSLDSWWTSLNFLGGTTVCLGQNSQSP-NH2
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           Ac-LELDKWASLWNWA-NH2
     609
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    743
          Ac-IVOOONNLLRAIEAOOHLLOLTVWGIKOLOARILAVER-NH2
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    745
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          Ac-YTSLIHSLIEESONOOEKNEOELLELDKWASLWNWFNI-NH2
    897
          AC-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWFNIT-NH2
    898
          Ac-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWFNITN-NH2
    899
          Ac-YDPLVFPSDEFDASISQVNEKINQSLAFIRKSDELLHNVNAGK-NH2
    900
          AC-NYTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWFN-NH2
          AC-NNYTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWFNI-NH2
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    905
          Ac-KCRAKFKOLLOHYREVAAAKSSENDRLRLLLKOMCPSLDVDSIIPRTPD-NH2
          AC-RAKFKOLLOHYREVAAAKSSENDRLRLLLKOMCPSLDVDSIIPRTPD-NH2
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    907
          AC-VYPSDEYDASISQVNEEINQALAYIAAADELLENV-NH2
    909
          Ac-YDASISQVNEEINQALAYIRKADELL-NH2
    910
          Ac-M-Nle-WMEWDREINNYTSLIHSLIEESQNQQEKNEQELLEL-NH2
          Ac-KNGTYDYPKYEEESKLNRNEIKGVKLSSMGVYQI-NH2
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    912
          Ac-VTEKIOMASDNINDLIQSGVNTRLLTIQSHVQNYI-NH2
    913
          QNQQEKNEQELLELDKWASLWNWF-NH2
    914
          AC-QNQQEKNEQELLELDKWASLWNWF-NH2
    915
          LWNWF-NH2
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    916
          ELLELDKWASLWNWF-NH2
    917
          EKNEQELLELDKWASLWNWF-NH2
          SLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
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    919
          AC-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNW
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          Ac-YTSLIHSLIEESQNQQEKNEQELLELDKWASLW
    921
     922
          Ac-YTSLIHSLIEESQNQQEKNEQELLELDKWASL
          TSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
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     924
          SLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
     925
          LIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
     926
          IHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
          Ac-AAVALLPAVLLALLAPSELEIKRYKNRVASRKCRAKFKQLLQHYREVAAAK-NH2
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     941
          Ac-AAVALLPAVLLALLAPCRAKFKQLLQHYREVAAAKSSENDRLRLLLKQMCP-NH2
     942
          Ac-YTSLIHSLIEESQNQQEKNNNIERDWEMWTMNNWIQ-NH2
     944
          VYPSDEYDASISQVNEEINQALAYIRKADELLENV-NH2
     945
          Ac-LMQLARQLMQLARQMKQLADSLMQLARQVSRLESA-NH2
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     946
          Ac-WMEWDREINNYTSLIHSLIEESQNQQEKNEQELL-NH2
     947
          Ac-MEWDREINNYTSLIHSLIEESQNQQEKNEQELLEL-NH2
     948
          Ac-EWDREINNYTSLIHSLIEESQNQQEKNEQELLEL-NH2
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    949
          Ac-MEWDREINNYTSLIHSLIEESQNQQEKNEQELLE-NH2
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          Biotin-W-Nle-EWDREINNYTSLIHSLIEESONOOEKNEOELLEL-NH2
    951
          AC-YLEYDREINNYTSLIHSLIEESQNQQEKNEQELLEL-NH2
    952
          Ac-IKQFINMWQEVGKAMYA-NH2
    953
          Ac-IRKSDELL-NH2
    954
          Decanoyl-IRKSDELL-NH2
    955
          Acetyl-Aca-Aca-IRKSDELL-NH2
    956
          Ac-YDASISOV-NH2
    957
          Ac-NEKINQSL-NH2
    958
          AC-SISQVNEEINQALAYIRKADELL-NH2
          Ac-QVNEEINQALAYIRKADELL-NH2
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    960
          Ac-EEINQALAYIRKADELL-NH
    961
          Ac-NQALAYIRKADELL-NH2
    962
          Ac-LAYIRKADELL-NH2
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    963
           FDASISQVNEKINQALAFIRKSDELL-NH2
    964
          Ac-W-Nle-EWDREINNYTSLIHSLIEESQNQQEKNEQELLEL-NH2
    965
          Ac-ASRKCRAKFKQLLQHYREVAAAKSSENDRLRLLLKQMCPSLDVDS-NH2
    967
          Ac-WLEWDREINNYTSLIHSLIEESQNQQEKNEQELLEL-NH2
    968
          AC-YVKGEPIINFYDPLVFPSDEFDASISQVNEKINQSL-NH2
    969
          Ac-VYPSDEYDASISQVNEEINQSLAYIRKADELLHNV-NH2
    970
          Ac-YDASISQVNEEINQALAYIRKADELLENV-NH2
    971
          Ac-YDASISQVNEEINQALAYIRKADELLE-NH2
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    972
          Ac-VYPSDEYDASISQVNEEINQALAYIRKAAELLHNV-NH2
    973
          Ac-VYPSDEYDASISQVNEEINQALAYIRKALELLHNV-NH2
    974
          Decanoyl-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
          Ac-VYPSDEYDASISQVNEEINQLLAYIRKLDELLENV-NH2
    975
          Ac-DEYDASISQVNEKINQSLAFIRKSDELL-NH2
    976
          Ac-SNDQGSGYAADKESTQKAFDGITNKVNSVIEKTNT-NH2
    977
    978
          Ac-ESTQKAFDGITNKVNSVIEKTNTQFEAVGKEFGNLEKR-NH2
    979
          Ac-DGITNKVNSVIEKTNTQFEAVGKEFGNLEKRLENLNK-NH2
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          Ac-DSNVKNLYDKVRSQLRDNVKELGNGAFEFYHK-NH2
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          Ac-RDNVKELGNGAFEFYHKADDEALNSVKNGTYDYPKY-NH2
    982
          AC-EFYHKADDEALNSVKNGTYDYPKY-NH2
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          AC-AAVALLPAVLLALLAPAADKESTQKAFDGITNKVNS-NH2
    984
          AC-AAVALLPAVLLALLAPAADSNVKNLYDKVRSQLRDN-NH2
    985
          Ac-KESTQKAFDGITNKVNSV-NH2
    986
          Ac-IEKTNTQFEAVGKEFGNLER-NH2
    987
          AC-RLENLNKRVEDGFLDVWTYNAELLVALENE-NH2
    988
          Ac-SNVKNLYDKVRSQLRDN-NH2
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          AC-WMEWDREINNYTSLIHSLIEESQNQQEKNEQEL-NH2
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    990
          AC-WMEWDREINNYTSLIHSLIEESQNQQEKNEQE-NH2
          Ac-MEWDREINNYTSLIHSLIEESQNQQEKNEQEL-NH2
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          Ac-MEWDREINNYTSLIHSLIEESQNQQEKNEQE-NH2
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    993
          Ac-EWDREINNYTSLIHSLIEESQNQQEKNEQELLE-NH2
    994
          AC-EWDREINNYTSLIHSLIEESQNQQEKNEQELL-NH2
    995
          Ac-EWDREINNYTSLIHSLIEESQNQQEKNEQEL-NH2
    996
          AC-YTKFIYTLLEESQNQQEKNEQELLELDKWASLWNWF-NH2
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    997
          Ac-YMKQLADSLMQLARQVSRLESA-NH2
          Ac-YLMQLARQMKQLADSLMQLARQVSRLESA-NH2
    998
          AC-YQEWERKVDFLEENITALLEEAQIQQEKNMYELQKL-NH2
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Sequence
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    1000 AC-WMAWAAAINNYTSLIHSLIEESONQOEKNEOEEEEE-NH2
    1001 AC-YASLIAALIEESQNQQEKNEQELLELAKWAALWAWF-NH2
    1002 [Ac-EWDREINNYTSLIHSLIEESQNQQEKNEQEGGC-NH2] dimer
    1003 Ac-YDISIELNKAKSDLEESKEWIKKSNOKLDSIGNWH-NH2
    1004 Biotinyl-IDISIELNKAKSDLEESKEWIKKSNQKLDSIGNWH-NH2
    1005 Ac-YTSLI-OH
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    1006 Fmoc-HSLIEE-OH
          Fmoc-SONOOEK-OH
    1007
    1008 Fmoc-NEOELLEL-OH
    1009 Fmoc-DKWASL-OH
    1010 Fmoc-WNWF-OH
    1011 AC-AKTLERTWDTLNHLLFISSALYKLNLKSVAQITLSI-NH2
    1012 Ac-NITLQAKIKQFINMWQEVGKAMYA-NH2
    1013 Ac-LENERTLDFHDSNVKNLYDKVRLOLRDN-NH2
    1014 Ac-LENERTLDFHDSNVKNLYDKVRLOLRDNVKELGNG-NH2
    1015 Ac-TLDFHDSNVKNLYDKVRLQLRDNVKELGNGAFEF-NH2
    1016 Ac-IDISIELNKAKSDLEESKEWIKKSNQKLDSIGNWH-NH2
          Biotinyl-SISQVNEEINQALAYIRKADELL-NH2
    1022 Biotinyl-SISQVNEEINQSLAYIRKSDELL-NH2
    1023 Ac-SISQVNEEINQSLAYIRKSDELL-NH2
    1024 Ac-IDISIELNKAKSDLEESKEWIEKSNQELDSIGNWE-NH2
    1025 Ac-IDISIELNKAKSDLEESKEWIKKSNQELDSIGNWH-NH2
    1026 Ac-IDISIELNKAKSDLEEAKEWIDDANQKLDSIGNWH-NH2
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    1027 Ac-IDISIELNKAKSDLEESKEWIKKANQKLDSIGNWH-NH2
    1028 Ac-IDISIELNKAKSDLEEAKEWIKKSNOKLDSIGNWH-NH2
    1029
          Biotinyl-NSVALDPIDISIELNKAKSDLEESKEWIKKSNQKL-NH2
    1030 Biotinyl-ALDPIDISIELNKAKSDLEESKEWIKKSNQKLDSI-NH2
    1031
          desAminoTyrosine-NSVALDPIDISIELNKAKSDLEESKEWIKKSNQKL-NH2
    1032
          desAminoTyrosine-ALDPIDISIELNKAKSDLEESKEWIKKSNQKLDSI-NH2
    1033
          Ac-YDASISOVNEEINOALAFIRKADEL-NH2
    1034 Ac-YDASISQVNEEINQSLAYIRKADELL-NH2
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    1035 Biotinyl-YDASISQVNEEINQALAYIRKADELL-NH2
    1036 Biotinyl-YDASISQVNEEINQSLAFIRKSDELL-NH2
    1037 Ac-YDASISQVNEEINQSLAFIRKSDELL-NH2
    1038 Ac-WLEWDREINNYTSLIHSLIEESONOOEKNEOEL-NH2
    1039 Biotinyl-IDISIELNKAKSDLEESKEWIRRSNQKLDSIGNWH-NH2
    1044 Ac-YESTQKAFDGITNKVNSVIEKTNTQFEAVGKEFGNLEKR-NH2
    1045 Biotin-DEYDASISQVNEKINQSLAFIRKSDELL-NH2
    1046 Ac-MEWDREINNYTSLIHSLIEESQNQQEKNEQELL-NH2
    1047
          Ac-WQEWEQKVRYLEANISQSLEQAQIQQEKNMYEL-NH2
    1048 Ac-WQEWEQKVRYLEANISOSLEOAOIOOEKNEYEL-NH2
    1049 Ac-WQEWEQKVRYLEANITALLEQAQIQQEKNEYEL-NH2
    1050 Ac-WQEWEQKVRYLEANITALLEQAQIQQEKNMYEL-NH2
    1051 Ac-WQEWEQKVRYLEANISQSLEQAQIQQEKNEYELQKL-NH2
    1052 Ac-WQEWEQKVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
    1053 Ac-WQEWEQKVRYLEANITALLEQAQIQQEKNMYELQKL-NH2
    1054 Ac-IDISIELNKAKSDLEESKEWIEKSNOKLDSIGNWH-NH2
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    1055 Ac-EFGNLEKRLENLNKRVEDGFLDVWTYNAELLVALENE-NH2
    1056 Ac-EDGFLDVWTYNAELLVLMENERTLDFHDSNVKNLYDKVRMQL-NH2
    1057 Ac-SISQVNEKINQSLAFIRKSDELL-NH2
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          Sequence
    No
          desaminoTyr-SISQVNEKINQSLAFIRKSDELL-NH2
    1058
          Ac-SISOVNEKINOSLAYIRKSDELL-NH2
    1060
          Ac-QQLLDVVKRQQEMLRLTVWGTKNLQARVTAIEKYLKDQ-NH2
    1061 YTSLIHSLIEESONOOEKNEOELLELDKWASLWNWFC
    1062 Ac-FDASISQVNEKINQSLAYIRKSDELL-NH2
    1063
          Ac-YTSLIHSLIEESQNQQEKNEQELLELDKWA
 5
          Indole-3-acetyl-DEFDASISQVNEKINQSLAFIRKSDELL-NH2
    1065
          Indole-3-acetyl-DEFDESISQVNEKINQSLAFIRKSDELL-NH2
    1066 Indole-3-acetyl-DEFDESISQVNEKIEQSLAFIRKSDELL-NH2
          Indole-3-acetyl-DEFDESISQVNEKIEESLAFIRKSDELL-NH2
    1067
          Indole-3-acetyl-DEFDESISQVNEKIEESLQFIRKSDELL-NH2
    1068
          Indole-3-acetyl-GGGGGDEFDASISQVNEKINQSLAFIRKSDELL-NH2
    1070
          2-Napthoyl-DEFDASISQVNEKINQSLAFIRKSDELL-NH2
          desnH2Tyr-DEFDASISQVNEKINQSLAFIRKSDELL-NH2
    1071
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    1072 biotin-ALDPIDISIELNKAKSDLEESKEWIRRSNQKLDSI-NH2
    1073
          Ac-YDASISQVNEKINQALAYIRKADELLHNVNAGKST-NH2
    1074
          Ac-VYPSDEYDASISQVNEKINQALAYIRKADELLHNV-NH2
          AC-VYPSDEYDASISOVNEKINOSLAYIRKSDELLHNV-NH2
    1075
          AC-WGWGYGYG-NH2
    1076
          Ac-YGWGWGWGF-NH2
    1077
    1078
          Ac-WQEWEQKVRYLEANITALQEQAQIQAEKAEYELQKL-NH2
          Ac-WQEWEQKVRYLEAEITALQEEAQIQAEKAEYELQKL-NH2
    1081 AC-YTSLIHSLIEESQNQQEKNEQELLELDKWAS
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    1082 AC-VWPSDEFDASISOVNEKINOSLAFIRKSDELLHNV-NH2
    1083
          Ac-SKNISEQIDQIKKDEQKEGTGWGLGGKWWTSDWGV-NH2
    1084
          Ac-LSKNISEQIDQIKKDEQKEGTGWGLGGKWWTSDWG-NH2
          Ac-DLSKNISEQIDQIKKDEQKEGTGWGLGGKWWTSDW-NH2
    1086
          Ac-EDLSKNISEQIDQIKKDEQKEGTGWGLGGKWWTSD-NH2
          Ac-IEDLSKNISEOIDOIKKDEOKEGTGWGLGGKWWTS-NH2
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    1088 Ac-GIEDLSKNISEQIDQIKKDEQKEGTGWGLGGKWWT-NH2
    1089 Ac-IGIEDLSKNISEQIDQIKKDEQKEGTGWGLGGKWW-NH2
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    1090 2-Napthoyl--PSDEFDASISQVNEKINQSLAFIRKSDELLHNVN-NH2
    1091 Ac-VYPSDEYDASISQVNEKINQALAYIRKADELLENV-NH2
    1092 Ac-VYPSDEFDASISQVNEKINQALAFIRKADELLENV-NH2
    1093 AC-VYPSDEYDASISQVNEKINOALAYIREADELLENV-NH2
          Biotinyl-YDASISQVNEKINQSLAFIRESDELL-NH2
    1094
          Ac-AIGIEDLSKNISEQIDQIKKDEQKEGTGWGLGGKW-NH2
    1095
    1096
          Ac-AAIGIEDLSKNISEQIDQIKKDEQKEGTGWGLGGK-NH2
    1097
          Ac-DAAIGIEDLSKNISEQIDQIKKDEQKEGTGWGLGG-NH2
    1098
          Ac-PDAAIGIEDLSKNISEQIDQIKKDEOKEGTGWGLG-NH2
    1099
          AC-NITDKIDQIIHDFVDKTLPDQGDNDNWWTGWRQWI-NH2
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          Ac-TKNITDKIDQIIHDFVDKTLPDQGDNDNWWTGWRQ-NH2
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          Ac-WTKNITDKIDQIIHDFVDKTLPDQGDNDNWWTGWR-NH2
          Ac-DWTKNITDKIDQIIHDFVDKTLPDQGDNDNWWTGW-NH2
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          AC-HDWTKNITDKIDQIIHDFVDKTLPDQGDNDNWWTG-NH2
          AC-PHDWTKNITDKIDQIIHDFVDKTLPDOGDNDNWWT-NH2
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    1106
           Ac-EPHDWTKNITDKIDQIIHDFVDKTLPDQGDNDNWW-NH2
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           Ac-IEPHDWTKNITDKIDQIIHDFVDKTLPDQGDNDNW-NH2
          Ac-Alephdwtknitdkidglihdfydktlpdggdndn-nh2
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          Ac-AAIEPHDWTKNITDKIDQIIHDFVDKTLPDQGDND-NH2
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          Ac-DAAIEPHDWTKNITDKIDOIIHDFVDKTLPDOGDN-NH2
    1110
    1111 Ac-LSPTVWLSVIWMMWYWGPSLYSILSPFLPLLPIFF-NH2
          Ac-GLSPTVWLSVIWMMWYWGPSLYSILSPFLPLLPIF-NH2
    1112
    1113
          Ac-VGLSPTVWLSVIWMMWYWGPSLYSILSPFLPLLPI-NH2
    1114
          Ac-FVGLSPTWLSVIWMMWYWGPSLYSILSPFLPLLP-NH2
    1115
          Ac-WFVGLSPTVWLSVIWMMWYWGPSLYSILSPFLPLL-NH2
    1116 Ac-OWFVFLSPTVWLSVIWMMWYWGPSLYSILSPFLPL-NH2
    1117
          Ac-VOWFVGLSPTVWLSVIWMMWYWGPSLYSILSPFLP-NH2
    1118 Ac-FVQWFVGLSPTVWLSVIWMMWYWGPSLYSILSPFL-NH2
          Ac-PFVQWFVGLSPTVWLSVIWMMWYWGPSLYSILSPF-NH2
    1119
          Ac-VPFVOWFVGLSPTVWLSVIWMMWYWGPSLYSILSP-NH2
    1120
          Ac-LVPFVQWFVGLSPTVWLSVIWMMWYWGPSLYSILS-NH2
    1121
          H-NHTTWMEWDREINNYTSLIHSLIEESQNQQEKNEQELLELDKW-OH
    1122
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    1123
          H-QARQLLSGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARILAVERYLKDQ-OH
          Ac-VYPSDEFDASISOVNEKINOSLAFIREADELLENV-NH2
    1124
    1125 Ac-VFPSDEFDASISQVNEKINQSLAYIREADELLENV-NH2
    1126 Ac-DEFDASISQVNEKINQSLAYIREADELL-NH2
          Ac-NEQELLELDKWASLWNWFGGGGDEFDASISQVNEKINQSLAFIRKSDELL-NH2
    1128 Ac-LELDKWASLWNWFGGGGDEFDASISOVNEKINOSLAFIRKSDELL-NH2
    1129 Naphthoyl-EGEGEGEGDEFDASISQVNEKINQSLAFIRKSDELL-NH2
    1130
          Ac-ASRKCRAKFKQLLQHYREVAAAKSSENDRLRLLLKQMCPSLDV-NH2
          Naphthoyl-GDEEDASISQVNEKINQSLAFIRKSDELL-NH2
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    1132 Naphthoyl-GDEEDASESQVNEKINQSLAFIRKSDELL-NH2
    1133 Naphthoyl-GDEEDASESQQNEKINQSLAFIRKSDELL-NH2
    1134 Naphthoyl-GDEEDASESQQNEKQNQSLAFIRKSDELL-NH2
    1135 Naphthoyl-GDEEDASESQQNEKQNQSEAFIRKSDELL-NH2
    1136 Ac-WGDEFDESISQVNEKIEESLAFIRKSDELL-NH2
    1137 AC-YTSLGGDEFDESISQVNEKIEESLAFIRKSDELLGGWNWF-NH2
          Ac-YTSLIHSLGGDEFDESISQVNEKIEESLAFIRKSDELLGGWASLWNWF-NH
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          2-Naphthoyl-GDEFDESISQVNEKIEESLAFIRKSDELL-NH2
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          2-Naphthov1-GDEEDESISOVOEKIEESLAFIRKSDELL-NH2
    1141
          2-Naphthoyl-GDEEDESISQVQEKIEESLLFIRKSDELL-NH2
    1142
          Biotin-GDEYDESISQVNEKIEESLAFIRKSDELL-NH2
    1143
          2-Naphthoyl-GDEYDESISQVNEKIEESLAFIRKSDELL-NH2
          AC-YTSLIHSLIDEQEKIEELAFIRKSDELLELDKWNWF-NH2
    1146 VYPSDEYDASISOVNEEINOALAYIRKADELLENV-NH2
    1147
          Ac-NNLLRAIEAQQHLLQLTVWGSKQLQARILAVERYLKDQ-NH2
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    1148
          GGGVYPSDEYDASISQVNEEINQALAYIRKADELLENV-NH2
    1149
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    1150
          Ac-PTRVNYILIIGVLVLAbuEVTGVRADVHLL-NH2
    1151
          Ac-PTRVNYILIIGVLVLAbuEVTGVRADVHLLEOPGNLW-NH2
          Ac-PEKTPLLPTRVNYILIIGVLVLAbuEVTGVRADVHLL-NH2
    1152
    1153 AhaGGGVYPSDEYDASISQVNEEINQALAYIRKADELLENV-NH2
    1155 Ac-YTSLIHSLGGDEFDESISQVNEKIEESLAFIRKSDELL-NH2
     1156 Ac-YTSLGGDEFDESISOVNEKIEESLAFIRKSDELL-NH2
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    1157
          Ac-DEFDESISQVNEKIEESLAFIRKSDELLGGWASLWNWF-NH2
    1158 Ac-DEFDESISQVNEKIEESLAFIRKSDELLGGWNWF-NH2
          Ac-YTSLIHSLIEESQNQQEKNEQELLELDKASLWNWF-NH2
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No.
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           AC-YTSLIHSLIEESQNQQEKNEQELLELDKSLWNWF-NH2
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     1162 AC-YTSLIHSLIEESQNQQEKNEQELLELDKWNWF-NH2
     1163 AC-MTWMEWDREINNYTSLIHSLIEESQNQQEKNEQELLELDKASLWNWF-NH2
     1164 AC-MTWMEWDREINNYTSLIHSLIEESQNQQEKNEQELLELDKSLWNWF-NH2
     1165 AC-MTWMEWDREINNYTSLIHSLIEESQNQQEKNEQELLELDKLWNWF-NH2
     1166
          AC-MTWMEWDREINNYTSLIHSLIEESQNQQEKNEQELLELDKWNWF-NH2
           Ac-mtwmewdreinnytslihslieesqnqqekneqelleldkwaslwn-nh2
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     1168
           Ac-mtwmewdreinnytslihslieesqnqqekneqelleldkwasl-nh2
     1169
           (Pyr) HWSY (2-napthyl-D-Ala) LRPG-NH2
     1170 Ac-WNWFDEFDESISQVNEKIEESLAFIRKSDELLWNWF-NH2
     1171 Ac-YTSLIHSLIEESQNQQEKNEQELLELDKYASLYNYF-NH2
     1172 AC-YTSLIHSLIEESQNQQEKNEQELLELDKYAYLYNYF-NH2
     1173 2-Naphthoyl-AcaAcaAcaDEFDESISQVNEKIEESLAFIRKSDELLAcaAcaAcaW-NH2
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     2-Naphthoyl-AcaAcaAcaGDEFDESISQVNEKIEESLAFIRKSDELLGAcaAcaAcaW-NH2
     1175
          2-Naphthoyl-GDEFDESISQVNEKIEESLAFIRESDELL-NH2
     1176 2-Naphthoyl-GDEFDESISQVNEKIEESLAFIEESDELL-NH2
     1177 Ac-WQEWEQKVNYLEANITALLEQAQIQQEKNEYELQKL-NH2
     1178 Ac-WQEWEQKVDYLEANITALLEQAQIQQEKNEYELQKL-NH2
     1179 Ac-WQEWEQKVRWLEANITALLEQAQIQQEKNEYELQKL-NH2
     1180 Ac-WQEWEKQVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
     1181 Ac-WQEWEHQVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
    1182 AC-WQEWEHKVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
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     1183 AC-WQEWDREVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
          Ac-WQEWEREVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
     1184
          Ac-WQEWERQVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
     1185
     1186 Ac-WQEWEQKVKYLEANITALLEQAQIQQEKNEYELQKL-NH2
    1187
          Ac-WQEWEQKVRFLEANITALLEQAQIQQEKNEYELOKL-NH2
    1188 Ac-VNalpsDEYDASISQVNEEINQALAYIRKADELLENV-NH2
    1189 Ac-VNalpSDEnalDASISQVNEEINQALAYIRKADELLENV-NH2
    1190 Ac-VNalPSDEYDASISQVNEEINQALANalIRKADELLENV-NH2
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    1191 Ac-VYPSDEFDASISQVNEKINQSLAFIREADELLFNFF-NH2
    1192
          Ac-VYPSDEYDASISQVNEEINOALAYIRKADELLFNFF-NH2
          AC-YTSLITALLEQAQIQQEKNEYELQKLDKWASLWNWF-NH2
    1193
          {\tt Ac-YTSLITALLEQAQIQQEKNEYELQKLDKWASLWEWF-NH2}
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    1195
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    1196 Ac-YTSLITALLEQAQIQQEKNEYELQELDEWASLWEWF-NH2
          AC-YTSLITALLEEAQIQQEKNEYELQELDEWASLWEWF-NH2
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          Naphthoyl-Aua-Aua-TALLEQAQIQQEKNEYELQKLAua-Aua-Aua-W-NH2
    1198
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          Ac-WQEAAQKVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
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          Ac-WQEWAAKVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
    1202
          AC-WQAAEQKVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
          Ac-WQEWEAAVRYLEANITALLEQAQIQQEKNEYELQKL-NH2
    1203
          Ac-wqeweqaaryleanitalleqaqiqqekneyelqkl-nh2
    1204
    1205
          Ac-woeweokaayleanitalleqaqiqqekneyelokl-nh2
    1206
          Ac-WQEWEQKVAALEANITALLEQAQIQQEKNEYELQKL-NH2
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    1207
          {\tt Ac-WQEWEQKVRYLEANITALLEQAQIQQEKNEYELQKLGGGGWASLWNF-NH2}
          2-Naphthoyl-GDEFDASISQVNEKINQSLAFIRKSDELT-NH2
    1208
          2-Naphthoyl-GDEFDASISQVNEKINQSLAFTRKSDELT-NH2
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          2-Naphthoy1-GDEFDASISQVNEKTNQSLAFTRKSDELT-NH2
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          2-Naphthoy1-GDEFDASISQTNEKTNQSLAFTRKSDELT-NH2
    1212 2-Naphthoyl-GDEFDASTSQTNEKTNQSLAFTRKSDELT-NH2
    1213 2-Naphthoyl-GDEYDASTSQTNEKTNQSLAFTRKSDELT-NH2
    1214 2-Naphthoyl-GDEFDEEISQVNEKIEESLAFIRKSDELL-NH2
    1215 2-Naphthoyl-GDEFDASISQVNEKINQSLAFIRKSDELA-NH2
    1216 2-Naphthoyl-GDEFDASASQANEKANQSLAFARKSDELA-NH2
    1217 2-Naphthoyl-GDEFDESISQVNEKIEESLAFTRKSDELL-NH2
    1218 2-Naphthoyl-GDEFDESISQVNEKTEESLAFIRKSDELL-NH2
    1219 2-Naphthoyl-GDEFDESISOTNEKIEESLAFIRKSDELL-NH2
    1220 2-Naphthoyl-GDEFDESTSOVNEKIEESLAFIRKSDELL-NH2
    1221 AC-WNWFDEFDESTSQVNEKIEESLAFIRKSDELLWNWF-NH2
    1222 AC-WNWFDEFDESTSQTNEKIEESLAFIRKSDELLWNWF-NH2
    1223 Ac-WNWFDEFDESTSQTNEKTEESLAFIRKSDELLWNWF-NH2
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    1224 Ac-LQAGFFLLTRILTIPQSLDSWWTSLNFLGGTTVAL-NH2
    1225 Ac-YTNLIYTLLEESQNQQEKNEQELLELDKWASLWSWF-NH2
    1226 Ac-WQEWEQKVRYLEANITALLEQAQIQQEKNEYELQKLDKWASLWNWF-NH2
    1227 AC-NNMTWQEWEQKVRYLEANITALLEQAQIQQEKNEYELQKLDKWASLWNWF-NH2
    1230 Ac-WNWFIEESDELLWNWF-NH2
    1231 2-Naphthcyl-GFIEESDELLW-NH2
    1232 Ac-WFIEESDELLW-NH2
    1233 2-Naphthoyl-GFNFFIEESDELLFNFF-NH2
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   1234 2-Naphthoyl-GESDELW-NH2
    1235 Ac-WNWFGDEFDESISQVQEEIEESLAFIEESDELLGGWNWF-NH2
    1236 AC-WNWFIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
    1237 AC-YTSLITALLEQAQIQQEENEYELQALDEWASLWEWF-NH2
    1238 AC-YTSLIHSLGGDEFDESISQVNEEIEESLAFIEESDELLGGWASLWNWF-NH2
          2-Naphthoyl-GDEFDESISQVQEEIEESLAFIEESDELL-NH2
    1240 H-QARQLLSSIMQQQNNLLRAIEAQQHLLQLTVWGIKQLQARILAVERYLKDQ-OH
    1241 Ac-CPKYVKQNTLKLATGMRNVPEKQTR-NH2
    1242 Ac-GLFGAIAGFIENGWEGMIDGWYGFRHONSC-NH2
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    1243 Ac-LNFLGGT-NH2
    1244 Ac-LDSWWTSLNFLGGT-NH2
    1245 Ac-ILTIPOSLDSWWTSLNFLGGT-NH2
    1246 Ac-GFFLLTRILTIPQSLDSWWTSLNFLGGT-NH2
    1247 Ac-WQEWEQKITALLEQAQIQQEKNEYELQKLDKWASLWNWF-NH2
          Ac-WNWFITALLEQAQIQQEKNEYELQKLDKWASLWNWF-NH2
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    1249 Ac-WQEWEQKITALLEQAQIQQEKNEYELQKLDKWASLWEWF-NH2
    1250 Ac-WQEWEQKVRYLEANITALLEQAQIQQEKIEYELQKL-NH2
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    1251 Ac-WQEWEQKVRYLEAQITALLEQAQIQQEKIEYELQKL-NH2
    1252 Ac-KENKANGTDAKVKLIKQELDKYKNAVTELQLLMQS-NH2
    1253 Ac-NIKENKANGTDAKVKLIKQELDKYKNAVTELQLLM-NH2
    1254
         (FS)-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
    1255 2-Naphthoyl-GWNWFAcaDEFDESISQVQEEIEESLAFIEESDELLAcaWNWF-NH2
    1256 AC-WNWFGDEFDESISQVNEKIEESLAFIEESDELLGWNWF-NH2
    1257
          Ac-WNWFGDEFDESISQVNEKIEESLAFIRKSDELLGWNWF-NH2
    1258 Ac-WNWF-Aca-DEFDESISQVNEKIEESLAFIRKSDELL-Aca-WNWF-NH2
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    1259 Ac-WNWF-Aca-DEFDESISQVNEKIEESLAFIEESDELL-Aca-WNWF-NH2
    1260 Ac-EESQNQQEKNEOELLELDKWA-NH2
    1261 EESQNQQEKNEQELLELDKWA
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No.
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    1262 AC-CGTTDRSGAPTYSWGANDTDVFVLNNTRPPLGNWFG-NH2
    1263 AC-GVEHRLEAACNWTRGERADLEDRDRSELSP-NH2
    1264 Ac-CVREGNASRAWVAVTPTVATRDGKLPT-NH2
    1265 Ac-CFSPRHHWTTQDANASIYPG-NH2
    1266 Ac-LQHYREVAAAKSSENDRLRLLLKQMCPSLDVDS-NH2
    1267
          AC-WOEWDREISNYTSLITALLEQAQIQQEKNEYELQKLDEWASLWEWF-NH2
          AC-CWQEWDREISNYTSLITALLEQAQIQQEKNEYELQKLDEWASLWEWFC-NH2
    1269
          Ac-WOEWDREISNYTSLITALLEOAOIOOEKNEYELOKLDEWEWF-NH2
    1270 Ac-CWQEWDREISNYTSLITALLEQAQIQQEKNEYELQKLDEWEWFC-NH2
    1271 Ac-GQNSQSPTSNHSPTSAPPTAPGYRWA-NH2
    1272 Ac-PGSSTTSTGPARTALTTAQGTSLYPSA-NH2
    1273 Ac-PGSSTTSTGPARTALTTAQGTSLYPSAAATKPSDGNATA-NH2
    1275 AC-WQEWDREITALLEQAQIQQEKNEYELQKLDKWASLWNWF-NH2
    1276 Ac-WQEWDREITALLEQAQIQQEKNEYELQKLDEWASLWEWF-NH2
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    1277
          Ac-WQEWDREITALLEQAQIQQEKNEYELQKLDEWEWF-NH2
    1278
          Ac-WQEWDREITALLEQAQIQQEKNEYELQKLDEWEWF-NH2
    1279
          Ac-WQEWEREITALLEQAQIQQEKNEYELQKLIEWEWF-NH2
    1280
          Ac-WQEWEREITALLEQAQIQQEKIEYELQKLDEWEWF-NH2
    1281 Ac-WQEWEITALLEQAQIQQEKNEYELQKLDEWEWF-NH2
    1282 Ac-WQEWEITALLEQAQIQQEKNEYELQKLIEWEWF-NH2
    1283 Ac-WQEWEITALLEQAQIQQEKIEYELQKLDEWEWF-NH2
    1284 Ac-WQEWEITALLEQAQIQQEKIEYELQKLIEWEWF-NH2
   1285 Ac-WOEWDREIDEYDASISOVNEKINOALAYIREADELWEWF-NH2
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    1286 Ac-WQEWEREIDEYDASISQVNEKINQALAYIREADELWEWF-NH2
          Ac-WQEWEIDEYDASISQVNEKINQALAYIREADELWEWF-NH2
    1287
    1288
          Ac-WQEWDREIDEYDASISQVNEEINQALAYIREADELWEWF-NH2
          Ac-WQEWEREIDEYDASISQVNEEINQALAYIREADELWEWF-NH2
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    1290
          AC-WQEWEIDEYDASISQVNEEINQALAYIREADELWEWF-NH2
          Ac-WQEWDEYDASISQVNEKINQALAYIREADELWEWF-NH2
    1291
    1292 Ac-WQEWDEYDASISQVNEEINQALAYIREADELWEWF-NH2
    1293 AC-WQEWEQKITALLEQAQIQQEKIEYELQKLIEWEWF-NH2
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    1294 Ac-WQEWEQKITALLEQAQIQQEKIEYELQKLIEWASLWEWF-NH2
    1295 Ac-WQEWEITALLEQAQIQQEKIEYELQKLIEWASLWEWF-NH2
    1298
          -VYPSDEYDASISQVNEEINQALAYIRKADELLENV-NH2
    1299 Ac-WVYPSDEYDASISQVNEEINQALAYIRKADELLENVWNWF-NH2
    1300 YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
    1301 Ac-WQEWDEYDASISQVNEKINQALAYIREADELWAWF-NH2
    1302 Ac-WQAWDEYDASISQVNEKINQALAYIREADELWAWF-NH2
          Ac-WQAWDEYDASISQVNEKINQALAYIREADELWEWF-NH2
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     1304 Biotin-YDPLVFPSDEFDASISQVNEKINOSLAFIRKSDEL-NH2
     1305 Biotin-YDPLVFPSDEFDASISQVNEKINQSLAF-NH2
     1306 Biotin-QVNEKINQSLAFIRKSDELLHNVNAGKST-NH2
     1307 Ac-WMEWDREI-NH2
     1308 Ac-WQEWEQKI-NH2
          Ac-WQEWEQKITALLEQAQIQQEKIEYELQKLIKWASLWEWF-NH2
     1310 Ac-WQEWEQKITALLEQAQIQQEKIEYELQKLIEWASLWEWF-NH2
    1311 Ac-WQEWEREISAYTSLITALLEQAQIQQEKIEYELQKLIEWEWF-NH2
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    1312
          Ac-WQEWEREISAYTSLITALLEQAQIQQEKIEYELQKEWEWF-NH2
    1313
          Ac-WQEWEREISAYTSLITALLEQAQIQQEKIEYELQKEWEW-NH2
          Ac-WQEWEREISAYTSLITALLEQAQIQQEKIEYELQKLIEWEW-NH2
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No.
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    1315
          Ac-FNLSDHSESIQKKFQLMKKHVNKIGVDSDPIGSWLR-NH2
    1316
          Ac-DHSESIQKKFQLMKKHVNKIGVDSDPIGSWLRGIF-NH2
    1317
          AC-WSVKQANLTTSLLGDLLDDVTSIRHAVLQNRA-NH2
    1318
          Biotin-WMEWDREI-NH2
    1319 Biotin-NNMTWMEWDREINNYTSL-NH2
    1320 Ac-GAASLTLTVQARQLLSGIVQQQNNLLRAIEAQQHLL-NH2
    1321 Ac-ASLTLTVQARQLLSGIVQQQNNLLRAIEAQQHLLQL-NH2
    1322 Ac-VSVGNTLYYVNKQEGKSLYVKGEPIINFYDPLVF-NH2
    1323 Ac-QHWSYGLRPG-NH2
    1324
          Ac-WQEWEQKIQHWSYGLRPGWASLWEWF-NH2
    1325
          Ac-WQEWEQKIQHWSYGLRPGWEWF-NH2
    1326 Ac-WNWFQHWSYGLRPGWNWF-NH2
    1327 Ac-FNFFQHWSYGLRPGFNFF-NH2
    1328 Ac-GAGAQHWSYGLRPGAGAG-NH2
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    1329 PLLVLQAGFFLLTRILTIPQSLDSWWTSLNFLGGT
    1330 Ac-WQEWEQKITALLEQAQIQQEKIEYELQKLAKWASLWEWF-NH2
    1331 Ac-WQEWEQKITALLEQAQIQQEKIEYELQKLAEWASLWEWF-NH2
    1332 AC-WQEWEQKITALLEQAQIQQEKAEYELQKLAEWASLWEWF-NH2
    1333
          AC-WQEWEQKITALLEQAQIQQEKAEYELOKLAEWASLWAWF-NH2
    1334
          AC-WQEWEQKITALLEQAQIQQEKAEYELQKLAKWASLWAWF-NH2
    1335
          Ac-TNKAVVSLSNGVSVLTSKVLDLKNYIDKQLLPIVNK-NH2
    1336 Ac-KAVVSLSNGVSVLTSKVLDLKNYIDKOLLPIVNKOS-NH2
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    1337 Ac-WQEWEQKITALLEQAQIQQEKNEYELQKLIEWEWF-NH2
    1338 Ac-WQEWEQKITALLEQAQIQQEKNEYELQKLIEWEWF-NH2
    1339 Ac-WQEWEQKITALLEQAQIQQEKIEYELQKLDKWEWF-NH2
    1340 Ac-YDPLVFPSDEFDASISQVNEKINQSLAF-NH2
    1341 Fluor--VYPSDEYDASISQVNEEINQALAYIRKADELLENV-NH2
          Fluor-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
          Ac-SGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARIL-NH2
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    1345 Ac-QQQNNLLRAIEAQQHLLQLTVWGIKQLQARILAVERYLKDQ-NH2
    1346 Ac-SGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARILAVERYLKDQ-NH2
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    1347 Ac-WQEWEQKITALLEQAQIQQEKNEYELQKLAEWASLWAWF-NH2
    1348 AC-WQEWEQKITALLEQAQIQQEKNEYELQKLAEWASLWAW-NH2
    1349 AC-WQEWEQKITALLEQAQIQQEKAEYELQKLAEWASLWAW-NH2
    1350 AC-WQEWEQKITALLEQAQIQQEKNEYELQKLAEWAGLWAWF-NH2
    1351 AC-WQEWEQKITALLEQAQIQQEKNEYELQKLAEWAGLWAW-NH2
    1352
          AC-WQEWEQKITALLEQAQIQQEKAEYELQKLAEWAGLWAW-NH2
          Ac-WQEWEQKITALLEQAQIQQEKNEYELQKLDKWAGLWEWF-NH2
    1353
          Ac-WQEWQHWSYGLRPGWEWF-NH2
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    1355 Ac-WQAWQHWSYGLRPGWAWF-NH2
    1356 Biotinyl-WQEWEQKITALLEQAQIQQEKNEYELQKLDKWASLWEWF-NH2
          WQEWEQKITALLEQAQIQQEKNEYELQKLDKWASLWEWF
    1358 WQEWEQKITALLEQAQIQQEKIEYELQKLIEWEWF
    1361 Ac-AGSTMGARSMTLTVQARQLLSGIVQQQNNLLRAIEAQQ-NH2
    1362 Ac-AGSAMGAASLTLSAQSRTLLAGIVQQQQQLLDVVKRQQ-NH2
    1363 AC-AGSAMGAASTALTAQSRTLLAGIVQQQQQLLDVVKRQQ-NH2
    1364 Ac-ALTAQSRTLLAGIVQQQQQLLDVVKRQQELLRLTVWGT-NH2
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    1365 Ac-TLSAQSRTLLAGIVQQQQQLLDVVKRQQEMLRLTVWGT-NH2
    1366 Ac-TLTVQARQLLSGIVQQQNNLLRAIEAQQHLLQLTVWGI-NH2
    1367 AC-WQAWIEYEAELSQVKEKIEOSLAYIREADELWAWF-NH2
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No.
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     1368
          Ac-WQAWIEYEASLSQAKEKIEESKAYIREADELWAWF-NH2
     1369 AC-WQAWIEYERLLVQAKLKIAIAKLYIAKELLEWAWF-NH2
     1370 AC-WQAWIEYERLLVQVKLKIAIALLYIAKELLEWAWF-NH2
    1371 AC-WQAWIELERLLVQVKLKLAIAKLEIAKELLEWAWF-NH2
    1372 Ac-GEWTYDDATKTFTVTEGGH-NH2
    1373 AC-WQEWEQKIGEWTYDDATKTFTVTEGGHWASLWEWF-NH2
    1374
          Ac-GEWTYDDATKTFTVTE-NH2
          AC-WQEWEQKIGEWTYDDATKTFTVTEWASLWEWF-NH2
    1376 Ac-MHRFDYRT-NH2
    1377 Ac-WQEWEQKIMHRFDYRTWASLWEWF-NH2
    1378 Ac-MHRFNWSTGGG-NH2
    1379 AC-WQEWEQKIMHRFNWSTGGGWASLWEWF-NH2
    1380 Ac-MHRFNWST-NH2
    1381 Ac-WQEWEQKIMHRFNWSTWASLWEWF-NH2
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    1382 Ac-LLVPLARIMTMSSVHGGG-NH2
          Ac-WQEWEQKILLVPLARIMTMSSVHGGGWASLWEWF-NH2
    1384 Ac-LLVPLARIMTMSSVH-NH2
    1385 Ac-WQEWEQKILLVPLARIMTMSSVHWASLWEWF-NH2
    1386 TALLEQAQIQQEKNEYELQKLDK
    1387 Ac-TALLEQAQIQQEKNEYELOKLDK-NH2
    1388 Ac-TALLEQAQIQQEKIEYELQKLIE-NH2
    1389
          TALLEQAQIQQEKIEYELQKLIE
    1390 Ac-QARQLLSGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARILAVERY-NH2
    1391 Rhod-QARQLLSGIVQQQNNLLRAIEAQQHLLQLTVWGIKQLQARILAVERY-NH2
          Ac-GAASLTLSAQSRTLLAGIVQQQQQLLDVVKRQQEML-NH2
    1393
          Ac-GSAMGAASLTLSAQSRTLLAGIVQQQQQLLDVVKRQQEML-NH2
    1394 Ac-PALSTGLIHLHQNIVDVQFLFGVGSSIASWAIKWEY-NH2
    1395 Ac-PALSTGLIHLHÓNIVDVQFLYGVGSSIASWAIK-NH2
    1396 AC-LSTTQWQVLPUSFTTLPALSTGLIHLHQNIVDVQY-NH2
    1397 AC-FRKFPEATFSRUGSGPRITPRUMVDFPFRLWHY-NH2
    1398 AC-DFPFRLWHFPUTINYTIFKVRLFVGGVEHRLEAAUNWTR-NH2&
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    1399 Ac-YVGGVEHRLEAAUNWTRGERUDLEDRDRSELSPL-NH2
    1400
          MVYPSDEYDASISQVNEEINQALAYIRKADELLENV
          Ac-GPLLVLQAGFFLLTRILTIPQSLDSWWTSLNFLGG-NH2
    1403
          Ac-LGPLLVLQAGFFLLTRILTIPQSLDSWWTSLNFLG-NH2
    1404 Ac-FLGPLLVLQAGFFLLTRILTIPQSLDSWWTSLNFL-NH2
    1405 AC-YTNTIYTLLEESQNQQEKNEQELLELDKWASLWNWF-NH2
    1406 YTNTIYTLLEESQNQQEKNEQELLELDKWASLWNWF
    1407 Ac-YTGIIYNLLEESQNQQEKNEQELLELDKWANLWNWF-NH2
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    1408 YTGIIYNLLEESQNQQEKNEQELLELDKWANLWNWF
    1409 Ac-YTSLIYSLLEKSQIQQEKNEQELLELDKWASLWNWF-NH2
    1410 YTSLIYSLLEKSQIQQEKNEQELLELDKWASLWNWF
          Ac-EKSQIQQEKNEQELLELDKWA-NH2
    1411
          EKSOIOOEKNEOELLELDKWA
    1413
          Ac-EQAQIQQEKNEYELQKLDKWA-NH2
    1414 AC-YTSLIGSLIEESQIQQERNEQELLELDRWASLWEWF-NH2
    1415 Ac-YTXLIHSLIXESQNQQXKNEQELXELDKWASLWNWF-NH2
    1416 Ac-YTXLIHSLIWESQNQQXKNEQELXELD-NH2
    1417 Ac-YTSLIHSLIEESQNQQEKNEQELLELD-NH2
    1418 Ac-WQEQEXKITALLXQAQIQQXKNEYELXKLDKWASLWEWF-NH2
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    No.
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    1419
          Ac-XKITALLXQAQIQQXKNEYELXKLDKWASLWEWF-NH2
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          AC-WEQKITALLEQAQIQQEKNEYELQKLD-NH2
    1421
          Ac-WEXKITALLXQAQIQQXKNEYELXKLD-NH2
    1422
          Ac-XKITALLXQAQIQQXKNEYELXKLD-NH2
    1425
          Ac-QKITALLEQAQIQQEKNEYELQKLD-NH2
    1426
          Ac-QKITALLEQAQIQQEKNEYELQKLDKWASLWEWF-NH2
    1427
          Ac-WQEWEQKITALLEQAQIQQEKNEYELQKLD-NH2
    1428
          AC-VYPSDEYDASISQVNEEINQALAYIRKADELLEN-OH
    1429
          AC-VYPSDEYDASISOVNEEINOALAYIRKADELLE-OH
          AC-VYPSDEYDASISQVNEEINQALAYIRKADELL-OH
    1430
    1431
          Ac-VYPSDEYDASISQVNEEINQALAYIRKADEL-OH
    1432
          YPSDEYDASISQVNEEINQALAYIRKADELLENV-NH2
    1433
          PSDEYDASISQVNEEINQALAYIRKADELLENV-NH2
          SDEYDASISOVNEEINOALAYIRKADELLENV-NH2
    1435
          DEYDASISQVNEEINQALAYIRKADELLENV-NH2
    1436
          Ac-VYPSDEYDASISQVDEEINQALAYIRKADELLENV-NH2
    1437
          Ac-VYPSDEYDASISQVNEEIDQALAYIRKADELLENV-NH2
    1438
          AC-VYPSDEYDASISOVNEEINOALAYIRKADELLEDV-NH2
    1439
          Ac-VYPSDEYDASISQVDEEIDQALAYIRKADELLENV-NH2
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          Ac-LLSTNKAVVSLSNGVSVLTSKVLDLKNYIDKQLLP-NH2
          Ac-LSTNKAVVSLSNGVSVLTSKVLDLKNYIDKQLLPI-NH2
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          AC-STNKAVVSLSNGVSVGTSKVLDLKNYIDKOLLPIV-NH2
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    1443
          Ac-TNKAVVSLSNGVSVLTSKVLDLKNYIDKQLLPIVN-NH2
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          Ac-NKAVVSLSNGVSVLTSKVLDLKNYIDKQLLPIVNK-NH2
          Ac-KAVVSLSNGVSVLTSKVLDLKNYIDKQLLPIVNKQ-NH2
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    1446
          Ac-AVVSLSNGVSVLTSKVLDLKNYIDKQLLPIVNKOS-NH2
          Ac-VVSLSNGVSVLTSKVDLKNYIDKQWLLPIVNKQSU-NH2
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    1448
          Ac-VSLSNGVSVLTSKVLDLKNYIDKQLLPIVNKQSUS-NH2
    1449
          Ac-SLSNGVSVLTSKVLDLKNYIDKQLLPIVNKQSUSI-NH2
    1450
          Ac-LSNGVSVLTSKVLDKLKNYIDKQLLPIVNKQSUSIS-NH2
          AC-SNGVSVLTSKVLDLKNYIDKQLLPIVNKQSUSISN-NH2
          Ac-NGVSVLTSKVLDLKNYIDKQLLPIVNKQSUSISNI-NH2
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          Ac-GVSVLTSKVLDLKNYIDKQLLPIVNKQSUSISNIE-NH2
          Ac-VSVLTSKVLDLKNYIDKQLLPIVNKQSUSISINIET-NH2
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          Ac-SVLTSKVLDLKNYIDKQLLPIVNKQSUSISNIETV-NH2
          Ac-VLTSKVLDLKNYIDKQLLPIVNKQSUSISNIETVI-NH2
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          Ac-LTSKVLDLKNYIDKQLLPIVNKQSUSISNIETVIE-NH2
    1457
    1458
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    1459
          Ac-SKVLDLKNYIDKQLLPIVNKQSUSISNIETVIEFO-NH2
          Ac-KVLDLKNYIDKQLLPIVNKQSUSISNIETVIEFQQ-NH2
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    1462
          Ac-LDLKNYIDKQLLPIVNKQSUSISNIETVIEFQQKN-NH2
    1463
          Ac-DLKNYIDKQLLPIVNKQSUSISNIETVIEFQQKNN-NH2
    1464
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    1465
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    1466
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    1467
          Ac-YIDKQLLPIVNKQSUSISNIETVIEFQQKNNRLLE-NH2
    1468
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          Ac-DKQLLPIVNKQSUSISNIETVIEFQQKNNRLLEIT-NH2
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T
    No.
          Sequence
          Ac-KQLLPIVNKQSUSISNIETVIEFQQKNNRLLEITR-NH2
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          QVNEEINQALAYIRKADELLENV-NH2
    1473
          VYPSDEYDASISQVNEEINQALAYIRKADELLENV
    1475 Ac-DEYDASISQVNEEINQALAYIREADEL-NH2
    1476 Ac-DEYDASISQVNEKINQALAYIREADEL-NH2
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    1479 Ac-YHKCDDECLNSVKNGTFDFPKFEEESKLNRNEIKGVKLSS-NH2
    1480 Ac-YHK-Abu-DDE-Abu-LNSVKNGTFDFPKFEEESKLNRNEIKGVKLSS-NH2
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    1491 AC-YTSLIHSLIEESQIQQEKNEYELLELDKWASLWEWF-NH2
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    1504
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    1510
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    1516 AC-WQEWEREIQQEKNEYELQKLDKWASLWEWF-NH2
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          Ac-WQEWQAQIQQEKGEYELQKLIEWEWF-NH2
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No.
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          PEG-YTSLITALLEQAQIQQERNEQELLELDEWASLWEWF-NH2
    1522
          AC-YTSLITALLEQAQIQQERNEQELLELDEWASLWEWF-NH2
    1523
         PEG-GWQEWEQRITALLEQAQIQQERNEYELQELDEWASLWEWF-NH2
    1526
    1527 AC-GWQEWEQRITALLEQAQIQQERNEYELQELDEWASLWEWF-NH2
    1528
         PEG-YTSLIGSLIEESQIQQERNEQELLELDRWASLWEWF-NH2
    1529
          PEG-GWQEWEQRITALLEQAQIQQERNEYELQRLDRWASLWEWF-NH2
    1530
          Ac-GWQEWEQRITALLEQAQIQQERNEYELQRLDRWASLWEWF-NH2
          PEG-GWOEWEQRITALLEOAOIOOERNEYELOELDRWASLWEWF-NH2
          Ac-GWQEWEQRITALLEQAQIQOERNEYELOELDRWASLWEWF-NH2
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    1534 Ac-YTSLIGSLIEESQNQQERNEQELLELDRWASLWNWF-NH2
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10
    1539 NEOELLELDK
    1540
          WASLWNWF-NH2
    1542
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    1548 Ac-WQEWEQKITALLEQAQIQAAANEYELOKLDKWASLWEWF-NH2
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          Ac-WQEWEQKITALLEQAQIQQEKNEYELQKLDAAASLWEWF-NH
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    1558 AC-ERTLDFHDS-NH2
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    1567
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    1568
          Ac-WNHGNITLGEWYNQTKDLQQKFYEIIMDIEQNNVQ-NH2
    1572
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    1573
         Ac-YTSLIHSLIEESQDQQEKNEQELLELDKWASLWNWF-NH2
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         Ac-YTSLIHSLIEESONOOEKNEOELLELDKWASLWNWF
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           Ac-YTSLIHSLIEESQNQQEKNEQELLELAKWASLWNWF-NH2
     1629
           AC-YTSLIHSLIEESQNQQEKAEQELLELDKWASLWNWF-NH2
     1630
     1631
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          Ac-WQEWEQKITALLEQAQIQQEKNAYELQKLDKWASLWEWF-NH2
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     1645
           Ac-EQELLELDK-NH2
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     1650
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     1652
     1653
           AC-YTSLIHSLIEESANQQEANEQELLELDKWASLWNWF-NH2
15
     1654
           AC-YTSLIHSLIEESQAQQEKNEQELLELDKWASLWNWF-NH2
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           AC-YTSLIHSAIEESQNQQEKNEQELLELDKWASLWNWF-NH2
     1657
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     1658
          AC-YTSLIHSLAEESQNQQEKNEQELLELDKWASLWNWF-NH2
     1659
          Ac-YTSAIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2
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     1661
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     1662
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     1663
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     1664
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     1665
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           AC-HTIDLTDSEMNKLFEKTRROLREN-NH2
     1667
     1668
           Ac-SEMNKLFEKTRRQLREN -NH2
     1669
           Ac-VFPSDEADASISQVNEKINQSLAFIRKSDELLHNV-NH2
     1670
           Ac-VFPSDEFAASISQVNEKINQSLAFIRKSDELLHNV-NH2
25
     1671
           Ac-VFPSDEFDASISAVNEKINQSLAFIRKSDELLHNV-NH2
     1672
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     1673
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     1677
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    1679
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     1680
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     1681
          Ac-VFPSDEFDASISQVNEKINQSLAAIRKSDELLHNV-NH2
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T No.	Sequence
1682	Ac-VFPSDEFDASISQVNEKINQSLAFIRKSDEALHNV-NH2
1683	Ac-VFPSDEFDASISQVNEKINQSLAFIRKSDELAHNV-NH2
1684	Ac-VFPSDEFDASISQVNEKINQSLAFIRKSDELLANV-NH2
1685	Ac-WQEWEQKITALLEQAQIQQAKNEYELQKLDKWASLWEWF-NH2
1687	Ac-wqeweqkitalleqaqiqqekneyelqaldkwaslwewf-nh2
1688	Ac-wqeweqkitalleqaqiqqekneyelqkadkwaslwewf-nh2

5.4. SYNTHESIS OF PEPTIDES

The peptides of the invention may be synthesized or prepared by techniques well known in the art. for example, Creighton, 1983, Proteins: Structures and Molecular Principles, W.H. Freeman and Co., NY, 5 which is incorporated herein by reference in its entirety. Short peptides, for example, can be synthesized on a solid support or in solution. peptides may be made using recombinant DNA techniques. Here, the nucleotide sequences encoding the peptides 10 of the invention may be synthesized, and/or cloned, and expressed according to techniques well known to those of ordinary skill in the art. See, for example, Sambrook, et al., 1989, Molecular Cloning, A Laboratory Manual, Vols. 1-3, Cold Spring Harbor Press, NY. 15

The peptides of the invention may alternatively be synthesized such that one or more of the bonds which link the amino acid residues of the peptides are non-peptide bonds. These alternative non-peptide bonds may be formed by utilizing reactions well known to those in the art, and may include, but are not limited to imino, ester, hydrazide, semicarbazide, and azo bonds, to name but a few. In yet another embodiment of the invention, peptides comprising the sequences described above may be synthesized with additional chemical groups present at their amino and/or carboxy termini, such that, for example, the stability, bioavailability, and/or inhibitory activity of the peptides is enhanced. For example, hydrophobic groups such as carbobenzoxyl, dansyl, or tbutyloxycarbonyl groups, may be added to the peptides' 30 amino termini. Likewise, an acetyl group or a 9fluorenylmethoxy-carbonyl group may be placed at the

peptides' amino termini. (See "X" Tables I to IV, above.) Additionally, the hydrophobic group, t-butyloxycarbonyl, or an amido group may be added to the peptides' carboxy termini. (See "Z" in Tables I to IV, above.)

Further, the peptides of the invention may be synthesized such that their steric configuration is altered. For example, the D-isomer of one or more of the amino acid residues of the peptide may be used, rather than the usual L-isomer.

10 Still further, at least one of the amino acid residues of the peptides of the invention may be substituted by one of the well known non-naturally occurring amino acid residues. Alterations such as these may serve to increase the stability, bioavailability and/or inhibitory action of the

peptides of the invention.

Any of the peptides described above may, additionally, have a macromolecular carrier group covalently attached to their amino and/or carboxy termini. Such macromolecular carrier groups may include, for example, lipid-fatty acid conjugates, polyethylene glycol, carbohydrates or additional peptides. "X", in Tables I to IV, above, may therefore additionally represent any of the above macromolecular carrier groups covalently attached to the amino terminus of a peptide, with an additional peptide group being preferred. Likewise, "Z", in Tables I to IV, may additionally represent any of the macromolecular carrier groups described above.

5.5. ASSAYS FOR ANTI-MEMBRANE FUSION ACTIVITY

Described herein, are methods for ability of a compound, such as the peptides of the invention, to

inhibit membrane fusion events. Specifically, assays for cell fusion events are described in Section 5.5.1, below, and assays for antiviral activity are described in Section 5.5.2, below.

5 5.5.1 ASSAYS FOR CELL FUSION EVENTS

Assays for cell fusion events are well known to those of skill in the art, and may be used in conjunction, for example, with the peptides of the invention to test the peptides' antifusogenic capabilities.

Cell fusion assays are generally performed in vitro. Such an assay may comprise culturing cells which, in the absence of any treatment would undergo an observable level of syncytial formation. For example, uninfected cells may be incubated in the presence of cells chronically infected with a virus that induces cell fusion. Such viruses may include, but are not limited to, HIV, SIV, or respiratory syncytial virus.

For the assay, cells are incubated in the presence of a peptide to be assayed. For each peptide, a range of peptide concentrations may be tested. This range should include a control culture wherein no peptide has been added.

Standard conditions for culturing cells, well
known to those of ordinary skill in the art, are used.
After incubation for an appropriate period (24 hours at 37°C, for example) the culture is examined microscopically for the presence of multinucleated giant cells, which are indicative of cell fusion and syncytial formation. Well known stains, such as crystal violet stain, may be used to facilitate the visualization of syncytial formation.

5.5.2 ASSAYS FOR ANTIVIRAL ACTIVITY

The antiviral activity exhibited by the peptides of the invention may be measured, for example, by easily performed in vitro assays, such as those described below, which can test the peptides' ability 5 to inhibit syncytia formation, or their ability to inhibit infection by cell-free virus. Using these assays, such parameters as the relative antiviral activity of the peptides, exhibit against a given strain of virus and/or the strain specific inhibitory 10 activity of the peptide can be determined.

A cell fusion assay may be utilized to test the peptides' ability to inhibit viral-induced, such as HIV-induced, syncytia formation in vitro. assay may comprise culturing uninfected cells in the presence of cells chronically infected with a syncytial-inducing virus and a peptide to be assayed. For each peptide, a range of peptide concentrations may be tested. This range should include a control culture wherein no peptide has been added. Standard conditions for culturing, well known to those of ordinary skill in the art, are used. After incubation for an appropriate period (24 hours at 37°C, for example) the culture is examined microscopically for the presence of multinucleated giant cells, which are indicative of cell fusion and syncytia formation.

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25 Well known stains, such as crystal violet stain, may be used to facilitate syncytial visualization. Taking HIV as an example, such an assay would comprise CD-4* cells (such as Molt or CEM cells, for example) cultured in the presence of chronically HIV-infected cells and a peptide to be assayed.

Other well known characteristics of viral infection may also be assayed to test a peptide's

antiviral capabilities. Once again taking HIV as an example, a reverse transcriptase (RT) assay may be utilized to test the peptides' ability to inhibit infection of CD-4 cells by cell-free HIV. Such an assay may comprise culturing an appropriate 5 concentration (<u>i.e.</u>, TCID₅₀) of virus and CD-4 cells in the presence of the peptide to be tested. Culture conditions well known to those in the art are used. As above, a range of peptide concentrations may be used, in addition to a control culture wherein no 10 peptide has been added. After incubation for an appropriate period (e.g., 7 days) of culturing, a cell-free supernatant is prepared, using standard procedures, and tested for the present of RT activity as a measure of successful infection. The RT activity may be tested using standard techniques such as those 15 described by, for example, Goff et al. (Goff, S. et al., 1981, J. Virol. 38:239-248) and/or Willey et al. (Willey, R. et al., 1988, J. Virol. 62:139-147). These references are incorporated herein by reference in their entirety.

Standard methods which are well-known to those of skill in the art may be utilized for assaying non-retroviral activity. See, for example, Pringle et al. (Pringle, C.R. et al., 1985, J. Medical Virology 17:377-386) for a discussion of respiratory syncytial virus and parainfluenza virus activity assay techniques. Further, see, for example, "Zinsser Microbiology", 1988, Joklik, W.K. et al., eds., Appleton & Lange, Norwalk, CT, 19th ed., for a general review of such techniques. These references are incorporated by reference herein in their entirety.

In addition, the Examples presented below, in Sections

> 17, 18, 26 and 27 each provide additional assays for the testing of a compound's antiviral capability.

In vivo assays may also be utilized to test, for example, the antiviral activity of the peptides of the invention. To test for anti-HIV activity, for 5 example, the in vivo model described in Barnett et al. (Barnett, S.W. et al., 1994, Science 266:642-646) may be used.

Additionally, anti-RSV activity can be assayed in vivo via well known mouse models. For example, RSV can be administered intranasally to mice of various inbred strains. Virus replicates in lungs of all strains, but the highest titers are obtained in P/N, C57L/N and DBA/2N mice. Infection of BALB/c mice produces an asymptomatic bronchiolitis characterized by lymphocytic infiltrates and pulmonary virus titers of 104 to 105 pfu/g of lung tissue (Taylor, G. et al., 1984, Infect. Immun. 43:649-655).

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Cotton rat models of RSV are also well known. Virus replicates to high titer in the nose and lungs of the cotton rat but produces few if any signs of inflammation.

5.6. USES OF THE PEPTIDES OF THE INVENTION

The peptides of the invention may be utilized as antifusogenic or antiviral compounds, or as compounds 25 which modulate intracellular processes involving coiled coil peptide structures. Further, such peptides may be used to identify agents which exhibit antifusogenic, antiviral or intracellular modulatory activity. Still further, the peptides of the invention may be utilized as organism or viral type/subtype-specific diagnostic tools.

The antifusogenic capability of the peptides of the invention may additionally be utilized to inhibit or treat/ameliorate symptoms caused by processes involving membrane fusion events. Such events may include, for example, virus transmission via cell-cell fusion, abnormal neurotransmitter exchange via cellfusion, and sperm-egg fusion. Further, the peptides of the invention may be used to inhibit free viral, such as retroviral, particularly HIV, transmission to uninfected cells wherein such viral infection involves 10 membrane fusion events or involves fusion of a viral structure with a cell membrane. Among the intracellular disorders involving coiled coil peptides structures which may be ameliorated by the peptides of the invention are disorders involving, for example, bacterial toxins.

With respect to antiviral activity, the viruses whose transmission may be inhibited by the peptides of the invention include, but are not limited to human retroviruses, such as HIV-1 and HIV-2 and the human T-lymphocyte viruses (HTLV-I and II), and non-human retroviruses such as bovine leukosis virus, feline sarcoma and leukemia viruses, simian immunodeficiency, sarcoma and leukemia viruses, and sheep progress pneumonia viruses.

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Non retroviral viruses whose transmission may be
inhibited by the peptides of the invention include,
but are not limited to human respiratory syncytial
virus, canine distemper virus, newcastle disease
virus, human parainfluenza virus, influenza viruses,
measles viruses, Epstein-Barr viruses, hepatitis B
viruses, and simian Mason-Pfizer viruses.

Non enveloped viruses whose transmission may be inhibited by the peptides of the invention include,

but are not limited to picornaviruses such as polio viruses, hepatitis A virus, enterovirus, echoviruses and coxsackie viruses, papovaviruses such as papilloma virus, parvoviruses, adenoviruses and reoviruses.

As discussed more fully, below, in Section 5.6.1 and in the Example presented, below, in Section 8, DP107, DP178, DP107 analog and DP178 analog peptides form non-covalent protein-protein interactions which are required for normal activity of the virus. Thus, the peptides of the invention may also be utilized as components in assays for the identification of compounds that interfere with such protein-protein interactions and may, therefore, act as antiviral agents. These assays are discussed, below, in Section 5.6.1.

As demonstrated in the Example presented below in 15 Section 6, the antiviral activity of the peptides of the invention may show a pronounced type and subtype specificity, i.e., specific peptides may be effective in inhibiting the activity of only specific viruses. This feature of the invention presents many advantages. One such advantage, for example, lies in the field of diagnostics, wherein one can use the antiviral specificity of the peptide of the invention to ascertain the identity of a viral isolate. With respect to HIV, one may easily determine whether a 25 viral isolate consists of an HIV-1 or HIV-2 strain. For example, uninfected CD-4 cells may be co-infected with an isolate which has been identified as containing HIV the DP178 (SEQ ID:1) peptide, after which the retroviral activity of cell supernatants may be assayed, using, for example, the techniques 30 described above in Section 5.2. Those isolates whose retroviral activity is completely or nearly completely

inhibited contain HIV-1. Those isolates whose viral activity is unchanged or only reduced by a small amount, may be considered to not contain HIV-1. Such an isolate may then be treated with one or more of the other DP178 peptides of the invention, and

5 subsequently be tested for its viral activity in order to determine the identify of the viral isolate. The DP107 and DP178 analogs of the invention may also be utilized in a diagnostic capacity specific to the type and subtype of virus or organism in which the specific peptide sequence is found. A diagnostic procedure as described, above, for DP178, may be used in conjunction with the DP107/DP178 analog of interest.

5.6.1. SCREENING ASSAYS

As demonstrated in the Example presented in 15 Section 8, below, DP107 and DP178 portions of the TM protein gp41, i.e., the HR1 and HR2 portions of gp41, respectively, form non-covalent protein-protein interactions. As is also demonstrated, the maintenance of such interactions is necessary for 20 normal viral infectivity. Thus, compounds which bind DP107; bind DP178, and/or act to disrupt normal DP107/DP178 protein-protein interactions may act as antifusogenic, antiviral or cellular modulatory agents. Described below are assays for the 25 identification of such compounds. Note that, while, for ease and clarity of discussion, DP107 and DP178 peptides will be used as components of the assays described, but it is to be understood that any of the DP107 analog or DP178 analog peptides described, above, in Sections 5.1 through 5.3 may also be 30 utilized as part of these screens for compounds.

For example, in certain embodiments the assays of the invention may be use DP107 and/or DP178 analogs that contain one or more amino acid residue truncations, deletions, insertions or substitutions. In particular, in one preferred embodiment, the DP107. 5 DP178, DP107-like and DP178-like peptides can comprise amino and/or carboxy-terminal insertions corresponding to about two to about fifty amino acids amino-to or carboxy-to the endogenous sequence from which the DP107, DP178, DP107-like or DP178-like peptide is derived. In another particular embodiment, the peptides used in the assays described herein further comprise additional, heterologous sequence useful for detecting, immobilizing and/or purifying the particular peptide. Such heterologous sequences include, but are not limited to maltose binding fusion 15 proteins containing a DP178, DP107, DP178-like or DP107-like sequence such as the M41 Δ 178 and MF5.1 maltose binding fusion proteins described in Sections 8 and 30, below.

reduced binding affinities and are therefore useful,
e.g., to screen for compounds which inhibit the
formation of or, alternatively, disrupt complexes
between DP107/DP178 complexes. Among such reduced
binding analogs are peptides exhibiting one or more
25 alanine insertion or substitutions, including, e.g.,
the peptides described in the examples presented in
Sections 30 and 31, below. It is understood that such
analogs which have reduced binding affinities,
including the analogs described in Sections 30 and 31
below, are also part of the present invention.

Compounds which may be tested for an ability to bind DP107, DP178, and/or disrupt DP107/DP178 interactions, and which therefore, potentially represent antifusogenic, antiviral or intracellular modulatory compounds, include, but are not limited to, peptides made of D- and/or L-configuration amino acids (in, for example, the form of random peptide libraries; see Lam, K.S. et al., 1991, Nature 354:82-84), phosphopeptides (in, for example, the form of random or partially degenerate, directed 10 phosphopeptide libraries; see, for example, Songyang, Z. et al., 1993, Cell 72:767-778), antibodies, and small organic or inorganic molecules. Synthetic compounds, natural products, and other sources of potentially effective materials may be screened in a variety of ways, as described in this Section. 15

Compounds that can be screened, tested and identified as modulating HR1/HR2, DP178/DP107 and/or DP178-like/DP107-like interactions utilizing the methods described herein can, in general, include, e.g., small molecules that are of a molecular weight up to about 1500 daltons. Test compounds, including small molecules, can include, but are not limited to, compounds obtained from any commercial source, including Aldrich (1001 West St. Paul Ave., Milwaukee, WI 53233), Sigma Chemical (P.O. Box 14508, St. Louis, MO 63178), Fluka Chemie AG (Industriestrasse 25, CH-9471 Buchs, Switzerland (Fluka Chemical Corp. 980 South 2nd Street, Ronkonkoma, NY 11779)), Eastman Chemical Company, Fine Chemicals (P.O Box 431, Kingsport, TN 37662), Boehringer Mannheim GmbH (Sandhofer Strasse 116, D-68298 Mannheim), Takasago (4 Volvo Drive, Rockleigh, NJ 07647), SST Corporation

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(635 Brighton Road, Clifton, NJ 07012), Ferro (111 West Irene Road, Zachary, LA 70791), Riedel-deHaen Aktiengesellschaft (P.O. Box D-30918, Seelze, Germany), PPG Industries Inc., Fine Chemicals (One PPG Place, 34th Floor, Pittsburgh, PA 15272). Further any kind of natural products may be screened using the methods of the invention, including microbial, fungal or plant extracts.

Furthermore, diversity libraries of test compounds, including small molecule test compounds, 10 may be commercially obtained from Specs and BioSpecs B.V. (Rijswijk, The Netherlands), Chembridge Corporation (San Diego, CA), Contract Service Company (Dolgoprudny, Moscow Region, Russia), Comgenex USA Inc. (Princeton, NJ), Maybridge Chemicals Ltd. 15 (Cornwall PL34 OHW, United Kingdom), and Asinex (Moscow, Russia). Combinatorial libraries of test compounds, including small molecule test compounds, can be may be generated as disclosed in Eichler & Houghten, 1995, Mol. Med. Today 1:174-180; Dolle, 20 1997, Mol. Divers. 2:223-236; Lam, 1997, Anticancer Drug Des. 12:145-167. These references are incorporated hereby by reference in their entirety. It is to be noted that such references also teach additional screening methods which may be employed for 25 the further testing of compounds identified via the methods of the invention and which can aid in identifying and isolating compounds which can

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represent leads and therapeutic compounds.

The compounds, antibodies, or other molecules identified may be tested, for example, for an ability to inhibit cell fusion or viral activity, utilizing, for example, assays such as those described, above, in Section 5.5.

5 Among the peptides which may be tested are soluble peptides comprising DP107 and/or DP178 domains, and peptides comprising DP107 and/or DP178 domains having one or more mutations within one or both of the domains, such as the M41-P peptide

10 described, below, in the Example presented in Section 8, which contains a isoleucine to proline mutation within the DP178 sequence.

In one embodiment of such screening methods is a method for identifying a compound to be tested for antiviral ability comprising:

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- 1 exposing at least one compound to a peptide comprising a DP107 peptide for a time sufficient to allow binding of the compound to the DP107 peptide;
 - 2 removing non-bound compounds; and
- 20 3 determining the presence of the compound bound to the DP107 peptide, thereby identifying an agent to be tested for

antiviral ability.

In a second embodiment of such screening methods 25 is a method for identifying a compound to be tested for antiviral ability comprising:

- (a) exposing at least one compound to a peptide comprising a DP178 peptide for a time sufficient to allow binding of the compound to the DP178 peptide;
 - (b) removing non-bound compounds; and

(c) determining the presence of the compound bound to the DP178 peptide, thereby identifying an agent to be tested for antiviral ability.

One method utilizing these types of approaches that may be pursued in the isolation of such DP107binding or DP178-binding compounds is an assay which would include the attachment of either the DP107 or the DP178 peptide to a solid matrix, such as, for example, agarose or plastic beads, microtiter plate 10 wells, petri dishes, or membranes composed of, for example, nylon or nitrocellulose. In such an assay system, either the DP107 or DP178 protein may be anchored onto a solid surface, and the compound, or test substance, which is not anchored, is labeled, either directly or indirectly (e.g., with a radioactive label such as 125I, an absorption label such as biotin, or a fluorescent label such as fluorescein or rhodamine). In practice, microtiter plates are conveniently utilized. The anchored component may be immobilized by non-covalent or covalent attachments. Non-covalent attachment may be accomplished simply by coating the solid surface with a solution of the protein and drying. Alternatively, an immobilized antibody, preferably a monoclonal antibody, specific for the protein may be used to 25 anchor the protein to the solid surface. The surfaces may be prepared in advance and stored.

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In order to conduct the assay, the labeled compound is added to the coated surface containing the anchored DP107 or DP178 peptide. After the reaction is complete, unreacted components are removed (e.g., by washing) under conditions such that any complexes

formed will remain immobilized on the solid surface.

The detection of complexes anchored on the solid surface can be accomplished in a number of ways.

Where the compound is pre-labeled, the detection of label immobilized on the surface indicates that complexes were formed. Where the labeled component is not pre-labeled, an indirect label can be used to detect complexes anchored on the surface; e.g., using a labeled antibody specific for the compound (the antibody, in turn, may be directly labeled or indirectly labeled with a labeled anti-Ig antibody).

Alternatively, such an assay can be conducted in a liquid phase, the reaction products separated from unreacted components, and complexes detected; e.g., using an immobilized antibody specific for DP107 or DP178, whichever is appropriate for the given assay, or ab antibody specific for the compound, i.e., the test substance, in order to anchor any complexes formed in solution, and a labeled antibody specific for the other member of the complex to detect anchored complexes.

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By utilizing procedures such as this, large numbers of types of molecules may be simultaneously screened for DP107 or DP178-binding capability, and thus potential antiviral activity.

Further, compounds may be screened for an ability
to inhibit the formation of or, alternatively, disrupt
DP107/DP178 complexes. Such compounds may then be
tested for antifusogenic, antiviral or intercellular
modulatory capability. For ease of description, DP107
and DP178 will be referred to as "binding partners."

Compounds that disrupt such interactions may exhibit
antiviral activity. Such compounds may include, but

are not limited to molecules such as antibodies, peptides, and the like described above.

The basic principle of the assay systems used to identify compounds that interfere with the interaction between the DP107 and DP178 peptides involves ⁵ preparing a reaction mixture containing peptides under conditions and for a time sufficient to allow the two peptides to interact and bind, thus forming a complex. In order to test a compound for disruptive activity, the reaction is conducted in the presence and absence 10 of the test compound, i.e., the test compound may be initially included in the reaction mixture, or added at a time subsequent to the addition of one of the binding partners; controls are incubated without the test compound or with a placebo. The formation of any complexes between the binding partners is then 15 detected. The formation of a complex in the control reaction, but not in the reaction mixture containing the test compound indicates that the compound interferes with the interaction of the DP107 and DP178 peptides.

The assay for compounds that interfere with the interaction of the binding partners can be conducted in a heterogeneous or homogeneous format.

Heterogeneous assays involve anchoring one of the binding partners onto a solid phase and detecting complexes anchored on the solid phase at the end of the reaction. In homogeneous assays, the entire reaction is carried out in a liquid phase. In either approach, the order of addition of reactants can be varied to obtain different information about the compounds being tested. For example, test compounds that interfere with the interaction between the binding partners, e.g., by competition, can be

identified by conducting the reaction in the presence of the test substance; i.e., by adding the test substance to the reaction mixture prior to or simultaneously with the binding partners. On the other hand, test compounds that disrupt preformed 5 complexes, e.g. compounds with higher binding constants that displace one of the binding partners from the complex, can be tested by adding the test compound to the reaction mixture after complexes have been formed. The various formats are described briefly below.

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In a heterogeneous assay system, one binding partner, e.g., either the DP107 or DP178 peptide, is anchored onto a solid surface, and its binding partner, which is not anchored, is labeled, either directly or indirectly (e.g., with a radioactive label such as 125I, an absorption label such as biotin, or a fluorescent label such as fluorescein or rhodamine). In practice, microtiter plates are conveniently utilized. The anchored species may be immobilized by non-covalent or covalent attachments. Non-covalent attachment may be accomplished simply by coating the solid surface with a solution of the protein and drying. Alternatively, an immobilized antibody specific for the protein may be used to anchor the protein to the solid surface. The surfaces may be 25 prepared in advance and stored.

In order to conduct the assay, the binding partner of the immobilized species is added to the coated surface with or without the test compound. After the reaction is complete, unreacted components are removed (e.g., by washing) and any complexes formed will remain immobilized on the solid surface.

The detection of complexes anchored on the solid surface can be accomplished in a number of ways.

Where the binding partner was pre-labeled, the detection of label immobilized on the surface indicates that complexes were formed. Where the binding partner is not pre-labeled, an indirect label can be used to detect complexes anchored on the surface; e.g., using a labeled antibody specific for the binding partner (the antibody, in turn, may be directly labeled or indirectly labeled with a labeled anti-Ig antibody). Depending upon the order of addition of reaction components, test compounds which inhibit complex formation or which disrupt preformed complexes can be detected.

Alternatively, the reaction can be conducted in a liquid phase in the presence or absence of the test compound, the reaction products separated from unreacted components, and complexes detected; e.g., using an immobilized antibody specific for one binding partner to anchor any complexes formed in solution, and a labeled antibody specific for the other binding partner to detect anchored complexes. Again, depending upon the order of addition of reactants to the liquid phase, test compounds which inhibit complex or which disrupt preformed complexes can be identified.

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In an alternate embodiment of the invention, a homogeneous assay can be used. In this approach, a preformed complex of the DP107 and DP178 peptides is prepared in which one of the binding partners is labeled, but the signal generated by the label is quenched due to complex formation (see, e.g., U.S. Patent No. 4,109,496 by Rubenstein which utilizes this approach for immunoassays). The addition of a test

substance that competes with and displaces one of the binding partners from the preformed complex will result in the generation of a signal above background. In this way, test substances which disrupt DP-107/ DP-178 protein-protein interaction can be identified.

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In still another embodiment of the invention. fluorescence polarization may be used in a homogenous assay. In this approach, complex formation is detected by measuring the polarization of a fluorescently labeled peptide (e.g., with fluorescein 10 or rhodamine) in a sample. Binding of the peptide to its complementary HR1 or HR2 binding domain in a larger molecular weight peptide or protein, such as in a maltose binding fusion protein described herein, alters the correlation time of the fluorescent moiety and thereby decreases the fluroescence polarization of the labeled peptide.

In an alternative screening assay, test compounds may be assayed for the their ability to disrupt a DP178/DP107 interaction, as measured immunometrically using an antibody specifically reactive to a DP107/DP178 complex (i.e., an antibody that recognizes neither DP107 nor DP178 individually). Such an assay acts as a competition assay, and is based on techniques well known to those of skill in the art.

The above competition assay may be described, by 25 way of example, and not by way of limitation, by using the DP178 and M41A178 peptides and by assaying test compounds for the disruption of the complexes formed by these two peptides by immunometrically visualizing DP178/M41A178 complexes via the human recombinant Fab, Fab-d, as described, below, in the Example presented in Section 8. M41A178 is a maltose binding fusion

protein containing a gp41 region having its DP178 domain deleted, and is described, below, in the Example presented in Section 8.

Utilizing such an assay, M41A178 may be immobilized onto solid supports such as microtiter ⁵ wells. A series of dilutions of a test compound may then be added to each M41A178-containing well in the presence of a constant concentration of DP-178 peptide. After incubation, at, for example, room temperature for one hour, unbound DP-178 and test 10 compound are removed from the wells and wells are then incubated with the DP178/M41A178-specific Fab-d antibody. After incubation and washing, unbound Fab-d is removed from the plates and bound Fab-d is quantitated. A no-inhibitor control should also be conducted. Test compounds showing an ability to 15 disrupt DP178/M41∆178 complex formation are identified by their concentration-dependent decrease in the level of Fab-d binding.

A variation of such an assay may be utilized to perform a rapid, high-throughput binding assay which is capable of directly measuring DP178 binding to M41\(\Delta\)178 for the determination of binding constants of the ligand of inhibitory constants for competitors of DP178 binding.

Such an assay takes advantage of accepted

25 radioligand and receptor binding principles. (See, for example, Yamamura, H.I. et al., 1985,

"Neurotransmitter Receptor Binding", 2nd ed., Raven Press, NY.) As above, M41\(\Delta\)178 is immobilized onto a solid support such as a microtiter well. DP178

binding to M41\(\Delta\)178 is then quantitated by measuring the fraction of DP178 that is bound as \(^{125}I-DP178\) and calculating the total amount bound using a value for

specific activity (dpm/µg peptide) determined for each labeled DP178 preparation. Specific binding to M41A178 is defined as the difference of the binding of the labeled DP178 preparation in the microtiter wells (totals) and the binding in identical wells containing, in addition, excess unlabeled DP178 (nonspecifics).

Because the binding affinity for native DP178 and DP107 is very high (including native DP178-like and DP107-like peptides from other species; e.g., 10 nM 10 for DP178 in HIV-1, and 2 nM for T112 in RSV), test compounds must exhibit high binding properties to interfere with or disrupt the DP178/DP107 binding interaction. Accordingly, in another non-limiting example of the above-described competitions assays, such assays can be performed using "modified" DP107 and/or DP178 peptides (e.g., DP107 and/or DP178 analogs) which have reduced binding affinities relatived to the unmodified "parent peptides". use of such modified DP107 and DP178 peptides greatly increases the sensitivity of the competition assays of the invention by identifying more compounds with inhibitory potential. The binding affinities of compounds identified in the assays can then be optimized, e.g., using standard medicinal chemistry techniques, to generate compounds that are more 25 powerful inhibitors of DP107/DP178 complex formation and are therefore useful, e.g., as antiviral reagents. Alternatively, compounds identified in the competition assays using DP107 and/or DP178 analogs with reduced binding affinities may, themselves, be useful, e.g., as antiviral reagents.

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The term "reduced affinity," as used herein, refers to a DP107, DP178, DP107-like or DP178-like peptide that interacts with and forms a DP107/DP178 peptide pair, a HR1/DP178 pair or an HR2/DP107 pair under competition assay conditions, but interacts with. 5 its "partner" to form such a pair with a lower affinity than would a DP107 or DP178 "parent" peptide from which the reduced affinity peptide is derived.

Generally, the binding affinity of a peptide can be expressed as a B₅₀ value, i.e., the concentration of 10 peptide necessary for 50% of the peptide molecules to bind to their target under a given set of conditions. Preferably, the B₅₀ value of a reduced affinity peptide will by at least twice, and more preferably at least five times, at least 10 times, at least 20 times, or at least 100 times the B_{50} value of the unmodified peptide from which it was derived.

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Modified DP107 and DP178 peptides that have reduced binding affinities may be generated according to any number of techniques that will be readily apparent to those skilled in the art. For example, in one embodiment modified DP107 and DP178 peptides with reduced binding affinities may be generated by generating truncated DP107 and DP178 peptides, respectively. Such peptides may be routinely synthesized and tested, e.g., by the above described 25 screening assays, to determine their binding affinities to their target. For example, as described in the example presented below in Section 30, reducing the length of the native RSV DP178-like peptide T112 from 35 to 28 amino acid residues resulted in a five fold drop in binding affinity (from 1 nM to 5 nM).

Generally, such truncation can be of 1, 2, 3, 4, 5, 6, 7, 8, 9 or 10 amino acid residues.

Alternatively, modified DP107 and DP178 peptides with reduced binding affinity may be identified and generated by identifying and substituting, inserting 5 or deleting amino acid residues. For example in one embodiment, which is also demonstrated in the example presented below in Section 30, modified DP107 and/or DP178 peptides may be routinely synthesized and assayed for reduced binding affinity by systematically 10 replacing one or more amino acid residues of the native DP107 or DP178 peptide with other amino acid residues and testing the binding affinity of the resulting peptide by techniques such as those described herein. Preferably, the substituted amino acid residues are neutral amino acid residues exhibiting relatively small side chains, such as. alanine or glycine.

Such substitutions can identify "key" amino acid residues and can be used in the competition assays of the invention. Alternatively, upon identification of key residues by such systematic substitutions, the key residues can be changed to other residues and the resulting, modified peptides can be tested for binding affinity.

Modified DP107 and/or DP178 peptides that have reduced binding affinities may still further be identified using principles of protein chemistry and design that are well known to those of skill in the art. Specifically, such principles may be used to identify those amino acid residues of a native DP107 or DP178 sequence that effect, e.g., solubility, binding affinity, or stability of the peptide. Thus,

for example, using known principles of amino acid chemistry and protein design one skilled in the art could identify amino acid residues in a native DP107 or DP178 peptide that affect the structure of the peptide.

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5.7 PHARMACEUTICAL FORMULATIONS, DOSAGES AND MODES OF ADMINISTRATION

The peptides of the invention may be administered using techniques well known to those in the art. Preferably, agents are formulated and administered systemically. Techniques for formulation and administration may be found in "Remington's Pharmaceutical Sciences", 18th ed., 1990, Mack Publishing Co., Easton, PA. Suitable routes may include oral, rectal, transmucosal, or intestinal administration; parenteral delivery, including intramuscular, subcutaneous, intramedullary injections, as well as, intrathecal, direct intraventricular, intravenous, intraperitoneal, intranasal, or intraocular injections, just to name a 20 few. For injection, the agents of the invention may be formulated in aqueous solutions, preferably in physiologically compatible buffers such as Hanks' solution, Ringer's solution, or physiological saline buffer. For such transmucosal administration. penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art.

In instances wherein intracellular administration of the peptides of the invention or other inhibitory agents is preferred, techniques well known to those of ordinary skill in the art may be utilized. For example, such agents may be encapsulated into

liposomes, then administered as described above. Liposomes are spherical lipid bilayers with aqueous interiors. All molecules present in an aqueous solution at the time of liposome formation are incorporated into the aqueous interior. The liposomal 5 contents are both protected from the external microenvironment and, because liposomes fuse with cell membranes, are effectively delivered into the cell cytoplasm. Additionally, due to their hydrophobicity, when small molecules are to be administered, direct intracellular administration may be achieved.

Nucleotide sequences encoding the peptides of the invention which are to be intracellularly administered may be expressed in cells of interest, using techniques well known to those of skill in the art. For example, expression vectors derived from viruses such as retroviruses, vaccinia viruses, adeno- associated viruses, herpes viruses, or bovine papilloma viruses, may be used for delivery and expression of such nucleotide sequences into the targeted cell population. Methods for the 20 construction of such vectors and expression constructs are well known. See, for example, Sambrook et al., 1989, Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Press, Cold Spring Harbor NY, and Ausubel et al., 1989, Current Protocols in Molecular 25 Biology, Greene Publishing Associates and Wiley Interscience, NY.

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With respect to HIV, peptides of the invention, particularly DP107 and DP178, may be used as therapeutics in the treatment of AIDS. In addition, the peptides may be used as prophylactic measures in previously uninfected individuals after acute exposure to an HIV virus. Examples of such prophylactic use of

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the peptides may include, but are not limited to, prevention of virus transmission from mother to infant and other settings where the likelihood of HIV transmission exists, such as, for example, accidents in health care settings wherein workers are exposed to 5 HIV-containing blood products. The successful use of such treatments do not rely upon the generation of a host immune response directed against such peptides.

Effective dosages of the peptides of the invention to be administered may be determined through 10 procedures well known to those in the art which address such parameters as biological half-life, bioavailability, and toxicity. Given the data presented below in Section 6, DP178, for example, may prove efficacious in vivo at doses required to achieve circulating levels of about 1 to about 10 ng per ml of peptide.

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A therapeutically effective dose refers to that amount of the compound sufficient to result in . amelioration of symptoms or a prolongation of survival in a patient. Toxicity and therapeutic efficacy of such compounds can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g., for determining the LDso (the dose lethal to 50% of the population) and the ED50 (the dose therapeutically effective in 50% of the 25 population). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio LD₅₀/ED₅₀. Compounds which exhibit large therapeutic indices are preferred. The data obtained from these cell culture assays and animal studies can be used in formulating a range of 30 dosage for use in humans. The dosage of such compounds lies preferably within a range of

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circulating concentrations that include the ED50 with little or no toxicity. The dosage may vary within this range depending upon the dosage form employed and the route of administration utilized. For any compound used in the method of the invention, the 5 therapeutically effective dose can be estimated initially from cell culture assays. A dose may be formulated in animal models to achieve a circulating plasma concentration range that includes the IC50 (e.g., the concentration of the test compound which 10 achieves a half-maximal inhibition of the fusogenic event, such as a half-maximal inhibition of viral infection relative to the amount of the event in the absence of the test compound) as determined in cell culture. Such information can be used to more accurately determine useful doses in humans. Levels in plasma may be measured, for example, by high performance liquid chromatography (HPLC).

The peptides of the invention may, further, serve the role of a prophylactic vaccine, wherein the host raises antibodies against the peptides of the 20 invention, which then serve to neutralize HIV viruses by, for example, inhibiting further HIV infection.

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Administration of the peptides of the invention as a prophylactic vaccine, therefore, would comprise administering to a host a concentration of peptides 25 effective in raising an immune response which is sufficient to neutralize HIV, by, for example, inhibiting HIV ability to infect cells. The exact concentration will depend upon the specific peptide to be administered, but may be determined by using standard techniques for assaying the development of an immune response which are well known to those of

ordinary skill in the art. The peptides to be used as vaccines are usually administered intramuscularly.

The peptides may be formulated with a suitable adjuvant in order to enhance the immunological response. Such adjuvants may include, but are not limited to mineral gels such as aluminum hydroxide; surface active substances such as lysolecithin, pluronic polyols, polyanions; other peptides; oil emulsions; and potentially useful human adjuvants such as BCG and Corynebacterium parvum. Many methods may be used to introduce the vaccine formulations described here. These methods include but are not limited to oral, intradermal, intramuscular, intraperitoneal, intravenous, subcutaneous, and intranasal routes.

Alternatively, an effective concentration of polyclonal or monoclonal antibodies raised against the peptides of the invention may be administered to a host so that no uninfected cells become infected by HIV. The exact concentration of such antibodies will vary according to each specific antibody preparation, but may be determined using standard techniques well known to those of ordinary skill in the art.

Administration of the antibodies may be accomplished using a variety of techniques, including, but not limited to those described in this section.

25 For all such treatments described above, the exact formulation, route of administration and dosage can be chosen by the individual physician in view of the patient's condition. (See e.g. Fingl et al., 1975, in "The Pharmacological Basis of Therapeutics", Ch. 1 pl).

It should be noted that the attending physician would know how to and when to terminate, interrupt, or

adjust administration due to toxicity, or to organ dysfunctions. Conversely, the attending physician would also know to adjust treatment to higher levels if the clinical response were not adequate (precluding toxicity). The magnitude of an administrated dose in the management of the oncogenic disorder of interest will vary with the severity of the condition to be treated and the route of administration. The dose and perhaps dose frequency, will also vary according to the age, body weight, and response of the individual patient. A program comparable to that discussed above may be used in veterinary medicine.

Use of pharmaceutically acceptable carriers to formulate the compounds herein disclosed for the practice of the invention into dosages suitable for systemic administration is within the scope of the 15 invention. With proper choice of carrier and suitable manufacturing practice, the compositions of the present invention, in particular, those formulated as solutions, may be administered parenterally, such as by intravenous injection. The compounds can be formulated readily using pharmaceutically acceptable carriers well known in the art into dosages suitable for oral administration. Such carriers enable the compounds of the invention to be formulated as tablets, pills, capsules, liquids, gels, syrups, 25 slurries, suspensions and the like, for oral ingestion by a patient to be treated.

Pharmaceutical compositions suitable for use in the present invention include compositions wherein the active ingredients are contained in an effective amount to achieve its intended purpose. Determination of the effective amounts is well within the capability

of those skilled in the art, especially in light of the detailed disclosure provided herein.

In addition to the active ingredients, these pharmaceutical compositions may contain suitable pharmaceutically acceptable carriers comprising

5 excipients and auxiliaries which facilitate processing of the active compounds into preparations which can be used pharmaceutically. The preparations formulated for oral administration may be in the form of tablets, dragees, capsules, or solutions.

The pharmaceutical compositions of the present invention may be manufactured in a manner that is itself known, <u>e.g.</u>, by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping or lyophilizing processes.

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15 Pharmaceutical formulations for parenteral · administration include aqueous solutions of the active compounds in water-soluble form. Additionally, suspensions of the active compounds may be prepared as appropriate oily injection suspensions. Suitable lipophilic solvents or vehicles include fatty oils such as sesame oil, or synthetic fatty acid esters, such as ethyl oleate or triglycerides, or liposomes. Aqueous injection suspensions may contain substances which increase the viscosity of the suspension, such 25 as sodium carboxymethyl cellulose, sorbitol, or dextran. Optionally, the suspension may also contain suitable stabilizers or agents which increase the solubility of the compounds to allow for the preparation of highly concentrated solutions.

Pharmaceutical preparations for oral use can be obtained by combining the active compounds with solid excipient, optionally grinding a resulting mixture,

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and processing the mixture of granules, after adding suitable auxiliaries, if desired, to obtain tablets or dragee cores. Suitable excipients are, in particular, fillers such as sugars, including lactose, sucrose, mannitol, or sorbitol; cellulose preparations such as, for example, maize starch, wheat starch, rice starch, potato starch, gelatin, gum tragacanth, methyl cellulose, hydroxypropylmethyl-cellulose, sodium carboxymethylcellulose, and/or polyvinylpyrrolidone (PVP). If desired, disintegrating agents may be added, such as the cross-linked polyvinyl pyrrolidone, 10 agar, or alginic acid or a salt thereof such as sodium alginate.

Dragee cores are provided with suitable coatings. For this purpose, concentrated sugar solutions may be used, which may optionally contain gum arabic, talc, polyvinyl pyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures. Dyestuffs or pigments may be added to the tablets or dragee coatings for identification or to characterize different combinations of active compound doses.

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Pharmaceutical preparations which can be used orally include push-fit capsules made of gelatin, as well as soft, sealed capsules made of gelatin and a plasticizer, such as glycerol or sorbitol. 25 push-fit capsules can contain the active ingredients in admixture with filler such as lactose, binders such as starches, and/or lubricants such as talc or magnesium stearate and, optionally, stabilizers. soft capsules, the active compounds may be dissolved or suspended in suitable liquids, such as fatty oils, liquid paraffin, or liquid polyethylene glycols. In addition, stabilizers may be added.

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6. EXAMPLE: DP178 (SEQ ID:1) IS A POTENT INHIBITOR OF HIV-1 INFECTION

In this example, DP178 (SEQ ID:1) is shown to be a potent inhibitor of HIV-1 mediated CD-4 cell-cell fusion and infection by cell free virus. 5 fusion assay, this peptide completely blocks virus induced syncytia formation at concentrations of from 1-10 ng/ml. In the infectivity assay the inhibitory concentration is somewhat higher, blocking infection at 90ng/ml. It is further shown that DP178 (SEQ ID:1) shows that the antiviral activity of DP178 (SEQ ID:1) is highly specific for HIV-1. Additionally, a synthetic peptide, DP-185 (SEQ ID:3), representing a HIV-1-derived DP178 homolog is also found to block HIV-1-mediated syncytia formation.

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6.1. MATERIALS AND METHODS

6.1.1. PEPTIDE SYNTHESIS

Peptides were synthesized using Fast Moc chemistry on an Applied Biosystems Model 431A peptide 20 synthesizer. Generally, unless otherwise noted, the peptides contained amidated carboxy termini and acetylated amino termini. Amidated peptides were prepared using Rink resin (Advanced Chemtech) while peptides containing free carboxy termini were synthesized on Wang (p-alkoxy-benzyl-alcohol) resin (Bachem). First residues were double coupled to the appropriate resin and subsequent residues were single coupled. Each coupling step was followed by acetic anhydride capping. Peptides were cleaved from the resin by treatment with trifluoracetic acid (TFA) (10ml), H₂O (0.5ml), thioanisole (0.5ml), ethanedithiol (0.25ml), and crystalline phenol (0.75g). Purifi-

cation was carried out by reverse phase HPLC.

Approximately 50mg samples of crude peptide were chromatographed on a Waters Delta Pak C18 column (19mm x 30cm, 15µ spherical) with a linear gradient;

H₂O/acetonitrile 0.1% TFA. Lyophilized peptides were stored desiccated and peptide solutions were made in water at about 1mg/ml. Electrospray mass spectrometry yielded the following results: DP178 (SEQ ID:1):4491.87 (calculated 4491.94); DP-180 (SEQ ID:2):4491.45 (calculated 4491.94); DP-185 (SEQ ID:3):not done (calculated 4546.97).

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6.1.2. VIRUS

The HIV-1 virus was obtained from R. Gallo (Popovic, M. et al., 1984, Science 224:497-508) and propagated in CEM cells cultured in RPMI 1640 5 containing 10% fetal calf serum. Supernatant from the infected CEM cells was passed through a 0.2 µm filter and the infectious titer estimated in a microinfectivity assay using the AA5 cell line to support virus replication. For this purpose, 25µl of 10 serial diluted virus was added to $75\mu l$ AA5 cells at a concentration of 2 x 105/ml in a 96-well microtitre plate. Each virus dilution was tested in triplicate. Cells were cultured for eight days by addition of fresh medium every other day. On day 8 post infection, supernatant samples were tested for virus 15 replication as evidenced by reverse transcriptase activity released to the supernatant. The TCID, was calculated according to the Reed and Muench formula (Reed, L.J. et al., 1938, Am. J. Hyg. 27:493-497). The titer of the HIV-1_{LAI} and HIV-1_{MN} stocks used for 20 these studies, as measured on the AA5 cell line, was approximately 1.4 x 106 and 3.8 x 104 TCID₅₀/ml, respectively.

6.1.3. CELL FUSION ASSAY

Approximately 7 x 10⁴ Molt cells were incubated

25 with 1 x 10⁴ CEM cells chronically infected with the

HIV-1_{LAI} virus in 96-well plates (one-half area cluster

plates; Costar, Cambridge, MA) in a final volume of

100µl culture medium as previously described

(Matthews, T.J. et al., 1987, Proc. Natl. Acad. Sci.

USA 84: 5424-5428). Peptide inhibitors were added in

a volume of 10µl and the cell mixtures were incubated

for 24 hr. at 37°C. At that time, multinucleated

giant cells were estimated by microscopic examination at a 40x magnification which allowed visualization of the entire well in a single field.

6.1.4. CELL FREE VIRUS INFECTION ASSAY

5 Synthetic peptides were incubated at 37°C with either 247 TCID₅₀ (for experiment depicted in FIG. 2), or 62 TCID₅₀ (for experiment depicted in FIG.3) units of HIV-1_{LAI} virus or 25 TCID₅₀ units of HIV-2_{NIHZ} and CEM CD4° cells at peptide concentrations of 0, 0.04, 0.4, 4.0, and 40μg/ml for 7 days. The resulting reverse transcriptase (RT) activity in counts per minute was determined using the assay described, below, in Section 6.1.5. See, Reed, L.J. et al., 1938, Am. J. Hyg. 27: 493-497 for an explanation of TCID₅₀ calculations.

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6.1.5. REVERSE TRANSCRIPTASE ASSAY

The micro-reverse transcriptase (RT) assay was adapted from Goff et al. (Goff, S. et al., 1981, J. Virol. 38:239-248) and Willey et al. (Willey, R. et 20 al., 1988, J. Virol. 62:139-147). Supernatants from virus/cell cultures are adjusted to 1% Triton-X100. A 10 μ l sample of supernatant was added to 50 μ l of RT cocktail in a 96-well U-bottom microtitre plate and the samples incubated at 37°C for 90 min. The RT 25 cocktail contained 75mM KCl, 2mM dithiothreitol, 5mM MgCl₂, 5μ g/ml poly A (Pharmacia, cat. No. 27-4110-01), 0.25 units/ml oligo dT (Pharmacia, cat. No. 27-7858-01), 0.05% NP40, 50mM Tris-HCl, pH 7.8, 0.5 μ M nonradioactive dTTP, and $10\mu\text{Ci/ml}$ $^{32}\text{P-dTTP}$ (Amersham, cat. No. PB.10167). 30

After the incubation period, $40\,\mu l$ of reaction mixture was applied to a Schleicher and Schuell (S+S)

NA45 membrane (or DE81 paper) saturated in 2 x SSC buffer (0.3M NaCl and 0.003M sodium citrate) held in a S+S Minifold over one sheet of GB003 (S+S) filter paper, with partial vacuum applied. Each well of the minifold was washed four times with 200µl 2xSSC, under full vacuum. The membrane was removed from the minifold and washed 2 more times in a pyrex dish with an excess of 2xSSC. Finally, the membrane was drained on absorbent paper, placed on Whatman #3 paper, covered with Saran wrap, and exposed to film overnight at -70°C.

6.2. RESULTS

6.2.1. PEPTIDE INHIBITION OF INFECTED CELL-INDUCED SYNCYTIA FORMATION

The initial screen for antiviral activity assayed peptides' ability to block syncytium formation induced by overnight co-cultivation of uninfected Molt4 cells with chronically HIV-1 infected CEM cells. The results of several such experiments are presented herein. In the first of these experiments, serial 20 DP178 (SEQ ID:1) peptide concentrations between $10\mu g/ml$ and 12.5ng/ml were tested for blockade of the cell fusion process. For these experiments, CEM cells chronically infected with either HIV-1LAI, HIV-1MN, HIV- 1_{RF} , or HIV- 1_{SF2} virus were cocultivated overnight with uninfected Molt 4 cells. The results (FIG. 4) show 25 that DP178 (SEQ ID:1) afforded complete protection against each of the HIV-1 isolates down to the lowest concentration of DP178 (SEQ ID:1) used. For HIVLAI inhibition, the lowest concentration tested was 12.5ng/ml; for all other HIV-1 viruses, the lowest 30 concentration of DP178 (SEQ ID:1) used in this study was 100ng/ml. A second peptide, DP-180 (SEQ ID:2),

containing the same amino acid residues as DP178 (SEQ ID:1) but arranged in a random order exhibited no evidence of anti-fusogenic activity even at the high concentration of 40µg/ml (FIG. 4). These observations indicate that the inhibitory effect of DP178 (SEQ ID:1) is primary sequence-specific and not related to non-specific peptide/protein interactions. The actual endpoint (i.e., the lowest effective inhibitory concentration) of DP178 inhibitory action is within the range of 1-10 ng/ml.

The next series of experiments involved the preparation and testing of a DP178 (SEQ ID:1) homolog for its ability to inhibit HIV-1-induced syncytia formation. As shown in FIG. 1, the sequence of DP-185 (SEQ ID:3) is slightly different from DP178 (SEQ ID:1) in that its primary sequence is taken from the HIV-1_{SF2} isolate and contains several amino acid differences relative to DP178 (SEQ ID:1) near the N terminus. As shown in FIG. 4, DP-185 (SEQ ID:3), exhibits inhibitory activity even at 312.5ng/ml, the lowest concentration tested.

The next series of experiments involved a comparison of DP178 (SEQ ID:1) HIV-1 and HIV-2 inhibitory activity. As shown in FIG. 5, DP178 (SEQ ID:1) blocked HIV-1-mediated syncytia formation at peptide concentrations below lng/ml. DP178 (SEQ ID:1) failed, however, to block HIV-2 mediated syncytia formation at concentrations as high as 10μg/ml. This striking 4 log selectivity of DP178 (SEQ ID:1) as an inhibitor of HIV-1-mediated cell fusion demonstrates an unexpected HIV-1 specificity in the action of DP178 (SEQ ID:1). DP178 (SEQ ID:1) inhibition of HIV-1-mediated cell fusion, but the peptide's inability to inhibit HIV-2 medicated cell fusion in the same cell

type at the concentrations tested provides further evidence for the high degree of selectivity associated with the antiviral action of DP178 (SEQ ID:1).

6.2.2. PEPTIDE INHIBITION OF INFECTION BY CELL-FREE VIRUS

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DP178 (SEQ ID:1) was next tested for its ability to block CD-4° CEM cell infection by cell free HIV-1 virus. The results, shown in FIG. 2, are from an experiment in which DP178 (SEQ ID:1) was assayed for its ability to block infection of CEM cells by an 10 HIV-1LAI isolate. Included in the experiment were three control peptides, DP-116 (SEQ ID:9), DP-125 (SEQ ID:8), and DP-118 (SEQ ID:10). DP-116 (SEQ ID:9) represents a peptide previously shown to be inactive. using this assay, and DP-125 (SEQ ID:8; Wild, C. et 15 al., 1992, Proc. Natl. Acad, Sci. USA 89:10,537) and DP-118 (SEQ ID:10) are peptides which have previously been shown to be active in this assay. Each concentration (0, 0.04, 0.4, 4, and $40\mu g/ml$) of peptide was incubated with 247 TCID, units of HIV-1, ar 20 virus and CEM cells. After 7 days of culture, cellfree supernatant was tested for the presence of RT activity as a measure of successful infection. The results, shown in FIG. 2, demonstrate that DP178 (SEQ ID:1) inhibited the de novo infection process mediated by the HIV-1 viral isolate at concentrations as low as 25 90ng/ml (IC50=90ng/ml). In contrast, the two positive control peptides, DP-125 (SEQ: ID:8) and DP-118 (SEQ ID:10), had over 60-fold higher IC50 concentrations of approximately $5\mu g/ml$.

In a separate experiment, the HIV-1 and HIV-2 inhibitory action of DP178 (SEQ ID:1) was tested with CEM cells and either HIV-1, or HIV-2, 62 TCID, 60

HIV-1_{LAI} or 25 GCID₅₀ HIV-2_{NIHZ} were used in these experiments, and were incubated for 7 days. As may be seen in FIG. 3, DP178 (SEQ ID:1) inhibited HIV-1 infection with an IC50 of about 31ng/ml. In contrast, DP178 (SEQ ID:1) exhibited a much higher IC50 for HIV-2_{NIHZ}, thus making DP178 (SEQ ID:1) two logs more potent as a HIV-1 inhibitor than a HIV-2 inhibitor. This finding is consistent with the results of the fusion inhibition assays described, above, in Section 6.2.1, and further supports a significant level of selectivity (<u>i.e.</u>, for HIV-1 over HIV-2).

7. EXAMPLE: THE HIV-1 INHIBITOR, DP178 (SEO ID:1) IS NON-CYTOTOXIC

In this Example, the 36 amino acid synthetic peptide inhibitor DP178 (SEQ ID:1) is shown to be noncytotoxic to cells in culture, even at the highest peptide concentrations $(40\mu g/ml)$ tested.

7.1. MATERIALS AND METHODS

Cell proliferation and toxicity assay:

20 Approximately 3.8x10⁵ CEM cells for each peptide concentration were incubated for 3 days at 37°C in T25 flasks. Peptides tested were DP178 (SEQ ID:1) and DP-116 (SEQ ID:9), as described in FIG. 1. Peptides were synthesized as described, above, in Section 6.1. The concentrations of each peptide used were 0, 2.5, 10, and 40μg/ml. Cell counts were taken at incubation times of 0, 24, 48, and 72 hours.

7.2. RESULTS

Whether the potent HIV-1 inhibitor DP178 (SEQ 30 ID:1) exhibited any cytotoxic effects was assessed by assaying the peptide's effects on the proliferation

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and viability of cells in culture. CEM cells were incubated in the presence of varying concentrations of DP178 (SEQ ID:1), and DP-116 (SEQ ID:9), a peptide previously shown to be ineffective as a HIV inhibitor (Wild, C. et al., 1992, Proc. Natl. Acad. Sci. USA 5 89:10,537-10,541). Additionally, cells were incubated in the absence of either peptide.

The results of the cytotoxicity study demonstrate that DP178 (SEQ ID:1) exhibits no cytotoxic effects on cells in culture. As can be seen, below, in Table VI, 10 even the proliferation and viability characteristics of cells cultured for 3 days in the presence of the highest concentration of DP178 (SEQ ID:1) tested $(40\mu g/ml)$ do not significantly differ from the DP-116 (SEQ ID:9) or the no-peptide controls. The cell proliferation data is also represented in graphic form in FIG. 6. As was demonstrated in the Working Example presented above in Section 6, DP178 (SEQ ID:1) completely inhibits HIV-1 mediated syncytia formation at peptide concentrations between 1 and 10ng/ml, and completely inhibits cell-free viral infection at concentrations of at least 90ng/ml. Thus, this study demonstrates that even at peptide concentrations greater than 3 log higher than the HIV inhibitory dose, DP178 (SEQ ID:1) exhibits no cytotoxic effects.

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Table VI

% Viability
at time (hours)

	Peptide								
5	<u>Peptide</u>	Concentration µg/ml	0	24	48	72			
	DP178 (SEQ ID:1)	40	98	97	95	97			
		10	98	97	98	98			
10		2.5	98	93	96	96			
		2.5	70	,,,		30			
	DP116 (SEQ ID:9)	40	98	95	98	97			
		10	98	95	93	98			
15		2.5	98	96	98	99			
	No Peptide	0	98	97	99	98			

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8. EXAMPLE: THE INTERACTION OF DP178 AND DP107

Soluble recombinant forms of gp41 used in the example described below provide evidence that the DP178 peptide associates with a distal site on gp41 whose interactive structure is influenced by the DP107 leucine zipper motif. A single mutation disrupting the coiled-coil structure of the leucine zipper domain transformed the soluble recombinant gp41 protein from an inactive to an active inhibitor of HIV-1 fusion.

30 This transformation may result from liberation of the potent DP178 domain from a molecular clasp with the

leucine zipper, DP107, determinant. The results also indicate that the anti-HIV activity of various gp41 derivatives (peptides and recombinant proteins) may be due to their ability to form complexes with viral gp41 and interfere with its fusogenic process.

5

8.1. MATERIALS AND METHODS

8.1.1. CONSTRUCTION OF FUSION PROTEINS AND GP41 MUTANTS

Construction of fusion proteins and mutants shown in FIG. 7 was accomplished as follows: the DNA sequence corresponding to the extracellular domain of gp41 (540-686) was cloned into the Xmn I site of the expression vector pMal-p2 (New England Biolab) to give M41. The gp41 sequence was amplified from pgtat

(Malim et al., 1988, Nature 355: 181-183) by using polymerase chain reaction (PCR) with upstream primer 5'-ATGACGCTGACGGTACAGGCC-3' (primer A) and downstream primer 5'-TGACTAAGCTTAATACCACAGCCAATTTGTTAT-3' (primer B). M41-P was constructed by using the T7-Gen

in vitro mutagenesis kit from United States Biochemicals (USB) following the supplier's

20 In vitro mutagenesis kit from United States
Biochemicals (USB) following the supplier's
instructions. The mutagenic primer (5'GGAGCTGCTTGGGGCCCCAGAC-3') introduces an Ile to Pro
mutation in M41 at position 578. M41\Delta107, from which
the DP-107 region has been deleted, was made using a
deletion mutagenic primer 5'-

CCAAATCCCCAGGAGCTGCTCGAGCTGCACTATACCAGAC-3' (primer C) following the USB T7-Gen mutagenesis protocol.
M41\Delta178, from which the DP-178 region has been deleted, was made by cloning the DNA fragment

30 corresponding to gp41 amino acids 540-642 into the

Xmn I site of pMal-p2. Primer A and 5'-ATAGCTTCTAGATTAATTGTTAATTTCTCTGTCCC-3' (primer D) were used in the PCR with the template pgtat to generate the inserted DNA fragments. M41-P was used as the template with primer A and D in PCR to generate M41-PΔ178. All inserted sequences and mutated residues were checked by restriction enzyme analysis and confirmed by DNA sequencing.

8.1.2. PURIFICATION AND CHARACTERIZATION OF FUSION PROTEINS

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The fusion proteins were purified according to the protocol described in the manufacturer's brochure of protein fusion and purification systems from New England Biolabs (NEB). Fusion proteins (10 ng) were analyzed by electrophoresis on 8% SDS polyacrylamide gels. Western blotting analysis was performed as described by Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual, 2d Ed, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, Ch. 18, pp. 64-75. An HIV-1 positive serum diluted 1000-fold, 20 or a human Fab derived from repertoire cloning was used to react with the fusion proteins. .The second antibody was HRP-conjugated goat antihuman Fab. An ECL Western blotting detection system (Amersham) was used to detect the bound antibody. A detailed protocol for this detection system was provided by the 25 manufacturer. Rainbow molecular weight markers (Amersham) were used to estimate the size of fusion proteins.

8.1.3. <u>CELL FUSION ASSAYS FOR ANTI-HIV ACTIVITY</u> 30 Cell fusion assays were performed as previously described (Matthews et al., 1987, Proc. Natl. Acad.

Sci. USA 84: 5424-5481). CEM cells (7 X 10°) were incubated with HIV-1_{IIIB} chronically infected CEM cells (10°) in 96-well flat-bottomed half-area plates (Costar) in 100 μ l culture medium. Peptide and fusion proteins at various concentrations in 10 μ l culture medium were incubated with the cell mixtures at 37°C for 24 hours. Multinucleated syncytia were estimated with microscopic examination. Both M41 and M41-P did not show cytotoxicity at the concentrations tested and shown in FIG. 8.

Inhibition of HIV-1 induced cell-cell fusion activity was carried out in the presence of 10 nM DP178 and various concentrations of M41Δ178 or M41-PΔ178 as indicated in FIG. 9. There was no observable syncytia in the presence of 10 nM DP178. No peptide or fusion protein was added in the control samples.

8.1.4. ELISA ANALYSIS OF DP178 BINDING TO THE LEUCINE ZIPPER MOTIF OF GP41

The amino acid sequence of DP178 used is: YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF. For enzyme 20 linked immunoassay (ELISA), M41Δ178 or M41-PΔ178 (5 μ g/ml) in 0.1M NaHCO₃, pH 8.6, were coated on 96 wells Linbro ELISA plates (Flow Lab, Inc.) overnight. Each well was washed three times with distilled water then blocked with 3% bovine serum albumin (BSA) for 2 hours. After blocking, peptides with 0.5% BSA in TBST 25 (40 mM Tris-HCl pH7.5, 150 mM NaCl, 0.05% Tween 20) were added to the ELISA plates and incubated at room temperature for 1 hour. After washing three times with TBST, Fab-d was added at a concentration of 10 ng/ml with 0.5% BSA in TBST. The plates were washed three times with TBST after incubation at room temperature for 1 hour. Horse radish peroxidase (HRP)

conjugated goat antihuman Fab antiserum at a 2000 fold dilution in TBST with 0.5% BSA was added to each well and incubated at room temperature for 45 minutes. The plates were then washed four times with TBST. The peroxidase substrate o-phenylene diamine (2.5 mg/ml) and 0.15% H₂O₂ were added to develop the color. The reaction was stopped with an equal volume of 4.5 N H₂SO₄ after incubation at room temperature for 10 minutes. The optical density of the stopped reaction mixture was measured with a micro plate reader

(Molecular Design) at 490 nm. Results are shown in FIG. 10.

8.2. RESULTS

8.2.1. THE EXPRESSION AND CHARACTERIZATION OF THE ECTODOMAIN OF gp41

15 As a step toward understanding the roles of the two helical regions in gp41 structure and function, the ectodomain of gp41 was expressed as a maltose binding fusion protein (M41) (FIG. 7). The fusogenic peptide sequence at the N-terminal of gp41 was omitted 20 from this recombinant protein and its derivatives to improve solubility. The maltose binding protein facilitated purification of the fusion proteins under relatively mild, non-denaturing conditions. Because the M41 soluble recombinant gp41 was not glycosylated, lacked several regions of the transmembrane protein 25 (i.e., the fusion peptide, the membrane spanning, and the cytoplasmic domains), and was expressed in the absence of gp120, it was not expected to precisely reflect the structure of native gp41 on HIV-1 virions. Nevertheless, purified M41 folded in a manner that preserved certain discontinuous epitopes as evidenced by reactivity with human monoclonal antibodies, 98-6,

126-6, and 50-69, previously shown to bind conformational epitopes on native gp41 expressed in eukaryotic cells (Xu et al., 1991, J. Virol. 65: 4832-4838; Chen, 1994, J. Virol. 68:2002-2010). Thus, at least certain regions of native gp41 defined by these 5 antibodies appear to be reproduced in the recombinant fusion protein M41. Furthermore, M41 reacted with a human recombinant Fab (Fab-d) that recognizes a conformational epitope on gp41 and binds HIV-1 virions as well as HIV-1 infected cells but not uninfected 10 cells as analyzed by FACS. Deletion of either helix motif, i.e., DP107 or DP178, of the M41 fusion protein eliminated reactivity with Fab-d. These results indicate that both helical regions, separated by 60 amino acids in the primary sequence, are required to maintain the Fab-d epitope.

ANTI-HIV ACTIVITY OF THE RECOMBINANT ECTODOMAIN OF GP41

The wild type M41 fusion protein was tested for anti-HIV-1 activity. As explained, supra, synthetic 20 peptides corresponding to the leucine zipper (DP107) and the C-terminal putative helix (DP178) show potent anti-HIV activity. Despite inclusion of both these regions, the recombinant M41 protein did not affect HIV-1 induced membrane fusion at concentrations as high as 50 μ M (Table VII, below).

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Table VII

DISRUPTION OF THE LEUCINE ZIPPER OF GP41 FREES THE ANTI-HIV MOTIF

5		DP107	<u>DP178</u>	<u>M41</u>	<u>M41-P</u>	<u>M41-PΔ178</u>
	Cell fusion (IC ₉₀) 1 μ M	1 nM >50 μM	83 nM	>50 μM		
	Fab-D binding (k _D)	-	-	3.5x10 ⁻⁹	2.5x10 ⁻⁸	
10	HIV infectiv- ity (IC ₉₀)	1 μM 80 nM	>16 µN	M 66 nM	>8 µM	ſ

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Antiviral Infectivity Assays. 20 μl of serially diluted virus stock was incubated for 60 minutes at ambient temperature with 20 μl of the indicated concentration of purified recombinant fusion protein in RPMI 1640 containing 10% fetal bovine serum and antibiotics in a 96-well microtiter plate. 20 μl of CEM4 cells at 6 x 10⁵ cells/ml were added to each well, and cultures were incubated at 37°C in a humidified CO₂ incubator. Cells were cultured for 9 days by the addition of fresh medium every 2 to 3 days. On days 5, 7, and 9 postinfection, supernatant samples were assayed for reverse transcriptase (RT) activity, as described below, to monitor viral replication. The 50% tissue culture infectious dose (TCID₅₀) was calculated for each condition according to the formula of Reed & Muench, 1937, Am. J. Hyg. 27:493-497. RT activity was determined by a modification of the published methods of Goff et al., 1981, J. Virol. 38:239-248 and Willey et al., 1988, J. Virol. 62:139-147 as described in Chen et al., 1993, AIDS Res. Human Retroviruses 9:1079-1086.

Surprisingly, a single amino acid substitution, proline in place of isoleucine in the middle of the leucine zipper motif, yielded a fusion protein (M41-P) which did exhibit antiviral activity (Table XXV and Fig. 8). As seen in Table XXV, M41-P blocked syncytia

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The affinity constants of Fab-d binding to the fusion proteins were determined using a protocol described by B. Friguet et al., 1985, J. Immunol. Method. 77:305-319.

^{- =} No detectable binding of Fab-d to the fusion proteins.

formation by 90% at approximately 85 nM and neutralized HIV-1,1118 infection by 90% at approximately 70 nM concentrations. The anti-HIV-1 activity of M41-P appeared to be mediated by the C-terminal helical sequence since deletion of that region from M41-P yielded an inactive fusion protein, M41-PΔ178 (Table XXV). This interpretation was reinforced by experiments demonstrating that a truncated fusion protein lacking the DP178 sequence, M41A178, abrogated the potent anti-fusion activity of the DP178 peptide 10 in a concentration-dependent manner (FIG. 9). same truncated fusion protein containing the proline mutation disrupting the leucine zipper, M41-PA178, was not active in similar competition experiments (FIG. 9). The results indicate that the DP178 peptide associates with a second site on gp41 whose 15 interactive structure is dependent on a wild type leucine zipper sequence. A similar interaction may occur within the wild type fusion protein, M41, and act to form an intramolecular clasp which sequesters the DP178 region, making it unavailable for anti-viral 20 activity.

A specific association between these two domains is also indicated by other human monoclonal Fab-d studies. For example, Fab-d failed to bind either the DP178 peptide or the fusion protein M41Δ178, but its epitope was reconstituted by simply mixing these two reagents together (FIG. 10). Again, the proline mutation in the leucine zipper domain of the fusion protein, M41-PΔ178, failed to reconstitute the epitope in similar mixing experiments.

9. EXAMPLE: METHOD FOR COMPUTER-ASSISTED IDENTIFICATION OF DP107-LIKE AND DP178-LIKE SEQUENCES

A number of known coiled-coil sequences have been well described in the literature and contain heptad 5 repeat positioning for each amino acid. Coiled-coil nomenclature labels each of seven amino acids of a heptad repeat A through G, with amino acids A and D tending to be hydrophobic positions. Amino acids E and G tend to be charged. These four positions (A, D, E, and G) form the amphipathic backbone structure of a 10 monomeric alpha-helix. The backbones of two or more amphipathic helices interact with each other to form di-, tri-, tetrameric, etc., coiled-coil structures. In order to begin to design computer search motifs, a series of well characterized coiled coils were chosen 15 including yeast transcription factor GCN4, Influenza Virus hemagglutinin loop 36, and human proto-oncogenes c-Myc, c-Fos, and c-Jun. For each peptide sequence, a strict homology for the A and D positions, and a list of the amino acids which could be excluded for the B, 20 C, E, F, and G positions (because they are not observed in these positions) was determined. Motifs were tailored to the DP107 and DP178 sequences by deducing the most likely possibilities for heptad positioning of the amino acids of HIV-1 Bru DP-107, which is known to have coiled-coil structure, and HIV-25 1 Bru DP178, which is still structurally undefined. The analysis of each of the sequences is contained in FIG. 12. For example, the motif for GCN4 was designed as follows:

1. The only amino acids (using standard single letter amino acid codes) found in the A or D positions of GCN4 were [LMNV].

All amino acids were found at B, C, E, F, and G
positions except {CFGIMPTW}.

3. The PESEARCH motif would, therefore, be written as follows:

```
[LMNV] - {CFGIMPTW} (2) - [LMNV] - {CFGIMPTW} (3) - [LMNV] - {CFGIMPTW} (2) - [LMNV] - {CFGIMPTW} (3) - [LMNV] - {CFGIMPTW} (2) - [LMNV] - {CFGIMPTW} (3) - [LMNV] - {CFGIMPTW} (2) - [LMNV] - {CFGIMPTW} (3)
```

Translating or reading the motif: "at the first A

position either L, M, N, or V must occur; at positions

B and C (the next two positions) accept everything

except C, F, G, I, M, P, T, or W; at the D position

either L, M, N, or V must occur; at positions E, F,

and G (the next 3 positions) accept everything except

C, F, G, I, M, P, T, or W." This statement is

contained four times in a 28-mer motif and five times

in a 35-mer motif. The basic motif key then would be:

[LMNV]-{CFGIMPTW}. The motif keys for the remaining

well described coiled-coil sequences are summarized in

FIG. 12.

The motif design for DP107 and DP178 was slightly different than the 28-mer model sequences described above due to the fact that heptad repeat positions are not defined and the peptides are both longer than 28 residues. FIG. 13 illustrates several possible sequence alignments for both DP107 and DP178 and also includes motif designs based on 28-mer, 35-mer, and full-length peptides. Notice that only slight differences occur in the motifs as the peptides are lengthened. Generally, lengthening the base peptide results in a less stringent motif. This is very useful in broadening the possibilities for identifying

DP107-or DP-178-like primary amino acid sequences referred to in this document as "hits".

In addition to making highly specific motifs for each type peptide sequence to be searched, it is also possible to make "hybrid" motifs. These motifs are 5 made by "crossing" two or more very stringent motifs to make a new search algorithm which will find not only both "parent" motif sequences but also any peptide sequences which have similarities to one, the other, or both "parents". For example, in FIG. 14 the 10 "parent" sequence of GCN4 is crossed with each of the possible "parent" motifs of DP-107. Now the hybrid motif must contain all of the amino acids found in the A and D positions of both parents, and exclude all of the amino acids not found in either parent at the other positions. The resulting hybrid from crossing 15 GCN4 or [LMNV] {CFGIMPTW} and DP107 (28-mer with the first L in the D position) or [ILQT] {CDFIMPST}, is [ILMNOTV] {CFIMPT}. Notice that now only two basic hybrid motifs exist which cover both framing possibilities, as well as all peptide lengths of the parent DP-107 molecule. FIG. 15 represents the "hybridizations" of GCN4 with DP-178. FIG. 16 represents the "hybridizations" of DP107 and DP178. It is important to keep in mind that the represented motifs, both parent and hybrid, are motif keys and not 25 the depiction of the full-length motif needed to actually do the computer search.

Hybridizations can be performed on any combination of two or more motifs. FIG. 17 summarizes several three-motif hybridizations including GCN4, DP107 (both frames), and DP178 (also both frames). Notice that the resulting motifs are now becoming much more similar to each other. In

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fact, the first and third hybrid motifs are actually subsets of the second and fourth hybrid motifs respectively. This means that the first and third hybrid motifs are slightly more stringent than the second and fourth. It should also be noted that with ' only minor changes in these four motifs, or by hybridizing them, a single motif could be obtained which would find all of the sequences. However, it should be remembered that stringency is also reduced. Finally, the most broad-spectrum and least-stringent 10 hybrid motif is described in FIG. 18 which summarizes the hybridization of GCN4, DP107 (both frames), DP178 (both frames), c-Fos, c-Jun, c-Myc, and Flu loop 36.

A special set of motifs was designed based on the fact that DP-178 is located only approximately ten amino acids upstream of the transmembrane spanning 15 region of gp4l and just C-terminal to a proline which separates DP107 and DP178. It has been postulated that DP178 may be an amphipathic helix when membrane associated, and that the proline might aid in the initiation of the helix formation. The same arrangement was observed in Respiratory Syncytial Virus; however, the DP178-like region in this virus also had a leucine zipper just C-terminal to the proline. Therefore, N-terminal proline-leucine zipper motifs were designed to analyze whether any other 25 viruses might contain this same pattern. are summarized in FIG. 19.

The PC/Gene protein database contains 5879 viral amino acid sequences (library file PVIRUSES; CD-ROM release 11.0). Of these, 1092 are viral enveloped or glycoprotein sequences (library file PVIRUSE1).

Tables V through XIV contain lists of protein sequence

names and motif hit locations for all the motifs searched.

10. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION
OF DP107 AND DP178-LIKE SEQUENCES
IN HUMAN IMMUNODEFICIENCY VIRUS

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FIG. 20 represents search results for HIV-1 BRU isolate gp41 (PC/Gene protein sequence PENV HV1BR). Notice that the hybrid motif which crosses DP-107 and DP-178 (named 107x178x4; the same motif as found in FIG. 16 found three hits including amino acids 550-10 599, 636-688, and 796-823. These areas include DP-107 plus eight N-terminal and four C-terminal amino acids; DP178 plus seven N-terminal and ten C-terminal amino acids; and an area inside the transmembrane region ... (cytoplasmic). FIG. 20 also contains the results obtained from searching with the motif named ALLMOTI5, for which the key is found in FIG. 17 ({CDGHP} {CFP}x5). This motif also found three hits including DP107 (amino acids 510-599), DP178 (615-717), and a cytoplasmic region (772-841). These hits overlap the 20 hits found by the motif 107x178x4 with considerable additional sequences on both the amino and carboxy termini. This is not surprising in that 107x178x4 is a subset of the ALLMOTI5 hybrid motif. Importantly, even though the stringency of ALLMOTI5 is considerably less than 107x178x4, it still selectively identifies 25 the DP107 and DP178 regions of gp41 shown to contain sequences for inhibitory peptides of HIV-1. The results of these two motif searches are summarized in Table V of U.S. Patent Application Serial No. 08/470,896 filed on June 6, 1995 (incorporated herein by reference in its entirety) under the PC/Gene protein sequence name PENV_HV1BR. The proline-leucine

zipper motifs also gave several hits in HIV-1 BRU including 503-525 which is at the very C-terminus of gp120, just upstream of the cleavage site (P7LZIPC and P12LZIPC); and 735-768 in the cytoplasmic domain of gp41 (P23LZIPC). These results are found in Tables

5 VIII, IX, and X under the same sequence name as mentioned above. Notice that the only area of HIV-1 BRU which is predicted by the Lupas algorithm to contain a coiled-coil region, is from amino acids 635-670. This begins eight amino acids N-terminal to the start and ends eight amino acids N-terminal to the end of DP178. DP107, despite the fact that it is a known coiled coil, is not predicted to contain a coiled-coil region using the Lupas method.

11. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP107-LIKE AND DP178-LIKE SEQUENCES IN HUMAN RESPIRATORY SYNCYTIAL VIRUS

FIG. 21 represents search results for Human Respiratory Syncytial Virus (RSV; Strain A2) fusion glycoprotein F1 (PC/Gene protein sequence name PVGLF 20 HRSVA). Motif 107x178x4 finds three hits including amino acids 152-202, 213-243, and 488-515. arrangement of these hits is similar to what is found in HIV-1 except that the motif finds two regions with similarities to DP-178, one just downstream of what would be called the DP107 region or amino acids 213-243, and one just upstream of the transmembrane region (also similar to DP178) or amino acids 488-515. Motif ALLMOTI5 also finds three areas including amino acids 116-202, 267-302, and 506-549. The proline-leucine 30 zipper motifs also gave several hits including amino acids 205-221 and 265-287 (P1LZIPC 265-280, P12LZIPC),

and 484-513 (P7LZIPC and P12LZIPC 484-506, P23LZIPC). Notice that the PLZIP motifs also identify regions which share location similarities with DP-178 of HIV-1.

12. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP107-LIKE AND DP178-LIKE SEQUENCES IN SIMIAN IMMUNODEFICIENCY VIRUS

Motif hits for Simian immunodeficiency Virus gp41 (AGM3 isolate; PC/Gene protein sequence name PENV SIVAG) are shown in FIG. 22. Motif 107x178x4 10 finds three hits including amino acids 566-593, 597-624, and 703-730. The first two hits only have three amino acids between them and could probably be combined into one hit from 566-624 which would represent a DP107-like hit. Amino acids 703 to 730 would then represent a DP178-like hit. ALLMOTI5 also finds three hits including amino acids 556-628 (DP107like), 651-699 (DP178-like), and 808-852 which represents the transmembrane spanning region. SIV also has one region from 655-692 with a high 20 propensity to form a coiled coil as predicted by the Lupas algorithm. Both 107x178x4 and ALLMOTI5 motifs find the same region. SIV does not have any PLZIP motif hits in gp41.

The identification of DP178/DP107 analogs for a second SIV isolate (MM251) is demonstrated in the Example presented, below, in Section 19.

- 13. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP107-LIKE AND DP178 LIKE SEQUENCES

 IN CANINE DISTEMPER VIRUS
- 30 Canine Distemper Virus (strain Onderstepoort)
 fusion glycoprotein F1 (PC/Gene Protein sequence name

PVGLF_CDVO) has regions similar to Human RSV which are predicted to be DP107-like and DP178-like (FIG. 23). Motif 107x178x4 highlights one area just C-terminal to the fusion peptide at amino acids 252-293. Amino acids 252-286 are also predicted to be coiled coil using the Lupas algorithm. Almost 100 amino acids Cterminal to the first region is a DP178-like area at residues 340-367. ALLMOTI5 highlights three areas of interest including: amino acids 228-297, which completely overlaps both the Lupas prediction and the 10 DP107-like 107x178x4 hit; residues 340-381, which overlaps the second 107x178x4 hit; and amino acids 568-602, which is DP178-like in that it is located just N-terminal to the transmembrane region. It also overlaps another region (residues 570-602) predicted by the Lupas method to have a high propensity to form 15 a coiled coil. Several PLZIP motifs successfully identified areas of interest including P6 and P12LZIPC which highlight residues 336-357 and 336-361 respectively; P1 and P12LZIPC which find residues 398-414; and P12 and P23LZIPC which find residues 562-589 and 562-592 respectively.

14. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP107-LIKE AND DP178-LIKE SEQUENCES IN NEWCASTLE DISEASE VIRUS

protein sequence name PVGLF_NDVA). Motif 107x178x4 finds two areas including a DP107-like hit at amino acids 151-178 and a DP178-like hit at residues 426-512. ALLMOTIS finds three areas including residues 117-182, 231-272, and 426-512. The hits from 426-512 include a region which is predicted by the Lupas

method to have a high coiled-coil propensity (460-503). The PLZIP motifs identify only one region of interest at amino acids 273-289 (Pl and 12LZIPC).

15. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP107-LIKE AND DP178-LIKE SEQUENCES IN HUMAN PARAINFLUENZA VIRUS

Both motifs 107x178x4 and ALLMOTI5 exhibit

DP107-like hits in the same region, 115-182 and 117
182 respectively, of Human Parainfluenza Virus (strain

NIH 47885; PC/Gene protein sequence name PVGLF_p13H4;

(FIG. 25). In addition, the two motifs have a DP178like hit just slightly C-terminal at amino acids 207241. Both motifs also have DP178-like hits nearer the
transmembrane region including amino acids 457-497 and
462-512 respectively. Several PLZIP motif hits are

also observed including 283-303 (P5LZIPC), 283-310

(P12LZIPC), 453-474 (P6LZIPC), and 453-481 (P23LZIPC).
The Lupas algorithm predicts that amino acids 122-176
may have a propensity to form a coiled-coil.

20 16. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP107-LIKE AND DP178-LIKE SEQUENCES OF INFLUENZA A VIRUS

FIG. 26 illustrates the Lupas prediction for a coiled coil in Influenza A Virus (strain A/Aichi/2/68) at residues 379-436, as well as the motif hits for 107x178x4 at amino acids 387-453, and for ALLMOTI5 at residues 380-456. Residues 383-471 (38-125 of HA2) were shown by Carr and Kim to be an extended coiled coil when under acidic pH (Carr and Kim, 1993, Cell 73: 823-832). The Lupas algorithm predicts a coiled-coil at residues 379-436. All three methods successfully predicted the region shown to actually

have coiled-coil structure; however, ALLMOTI5 predicted the greatest portion of the 88 residue stretch.

17. EXAMPLE: POTENTIAL RESPIRATORY SYNCYTIAL VIRUS
DP178/DP107 ANALOGS: CD AND
ANTIVIRAL CHARACTERIZATION

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In the Example presented herein, respiratory syncytial virus (RSV) peptides identified by utilizing the computer-assisted search motifs described in the Examples presented in Sections 9 and 11, above, were tested for anti-RSV activity. Additionally, circular dichroism (CD) structural analyses were conducted on the peptides, as discussed below. It is demonstrated that several of the identified peptides exhibit potent antiviral capability. Additionally, it is shown that several of these peptides exhibit a substantial helical character.

17.1 MATERIALS AND METHODS

Structural analyses: The CD spectra were

measured in a 10mM sodium phosphate, 150mM sodium

chloride, pH 7.0, buffer at approximately 10mM

concentrations, using a 1 cm pathlength cell on a

Jobin/Yvon Autodichrograph Mark V CD

spectrophotometer. Peptides were synthesized

according to the methods described, above, in Section

6.1. Peptide concentrations were determined from A₂₈₀

using Edlehoch's method (1967, Biochemistry 6:1948).

Anti-RSV antiviral activity assays: The assay utilized herein tested the ability of the peptides to disrupt the ability of HEp2 cells acutely infected with RSV (i.e., cells which are infected with a multiplicity of infection of greater than 2) to fuse

and cause syncytial formation on a monolayer of uninfected an uninfected line of Hep-2 cells. The lower the observed level of fusion, the greater the antiviral activity of the peptide was determined to be.

Uninfected confluent monolayers of Hep-2 cells were grown in microtiter wells in 3% EMEM (Eagle Minimum Essential Medium w/o L-glutamine [Bio Whittaker Cat. No. 12-125F], with fetal bovine serum [FBS; which had been heat inactivated for 30 minutes at 56°C; Bio Whittaker Cat. No. 14-501F) supplemented at 3%, antibiotics (penicillin/streptomycin; Bio Whittaker Cat. No. 17-602E) added at 1%, and glutamine added at 1%.

To prepare Hep2 cells for addition to uninfected cells, cultures of acutely infected Hep2 cells were washed with DPBS (Dulbecco's Phosphate Buffered Saline w/o calcium or magnesium; Bio Whittaker Cat. No. 17-512F) and cell monolayers were removed with Versene (1:5000; Gibco Life Technologies Cat. No. 15040-017). The cells were spun 10 minutes and resuspended in 3% FBS. Cell counts were performed using a hemacytometer. Persistent cells were added to the uninfected Hep-2 cells.

The antiviral assay was conducted by, first, removing all media from the wells containing
25 uninfected Hep-2 cells, then adding peptides (at the dilutions described below) in 3% EMEM, and 100 acutely RSV-infected Hep2 cells per well. Wells were then incubated at 37°C for 48 hours.

After incubation, cells in control wells were checked for fusion centers, media was removed from the wells, followed by addition, to each well, of either Crystal Violet stain or XTT. With respect to Crystal

Violet, approximately $50\mu l$ 0.25% Crystal Violet stain in methanol were added to each well. The wells were rinsed immediately, to remove excess stain, and were allowed to dry. The number of syncytia per well were then counted, using a dissecting microscope.

With respect to XTT (2,3-bis[2-Methoxy-4-nitro-5-sulfophenyl]-2H-tetrazolium-5-carboxyanilide inner salt), 50μl XTT (1mg/ml in RPMI buffered with 100mM HEPES, pH 7.2-7.4, plus 5% DMSO) were added to each well. The OD_{450/690} was measured (after blanking against growth medium without cells or reagents, and against reagents) according to standard procedures.

<u>Peptides</u>: The peptides characterized in the study presented herein were:

- peptides T-142 to T-155 and T-575, as shown in FIG.
 27A, and peptides T-22 to T-27, T-68, T-334 and T-371 to T-375 and T-575, as shown in FIG. 27B;
 peptides T-120 to T-141 and T-576, as shown in FIG. 27B, and peptides T-12, T-13, T-15, T-19, T-28 to T-30, T-66, T-69, T-70 and T-576, as shown in FIG. 27D; and
- 3) peptides T-67 and T-104 to T-119 and T-384, as shown in FIG. 28A, and peptides T-71, T-613 to T-617, T-662 to T-676 and T-730, as shown in FIG. 28B.

The peptides of group 1 represent portions of the RSV F2 protein DP178/107-like region. The peptides of group 2 represent portions of the RSV F1 protein DP107-like region. The peptides of groups 3 represent portions of the RSV F1 protein DP178-like region.

Each peptide was tested at 2-fold serial dilutions ranging from $100\mu g/ml$ to approximately 100ng/ml. For each of the assays, a well containing no peptide was also used. The IC_{50} data for each

peptide represents the average of several experiments conducted utilizing that peptide.

17.2 RESULTS

The data summarized in FIGS. 27A-B and 28A-B

represent antiviral and structural information obtained from peptides derived from the RSV F2

DP178/DP107-like F2 region (FIG. 27A-B), the RSV F1

DP-107-like region (FIG. 27C-D) and the RSV DP178-like F2 region (FIG. 28A-B).

As shown in FIGS. 27A-D, a number of the RSV 10 DP178/DP107-like peptides exhibited a detectable level of antiviral activity. Peptides from the RSV DP178/DP107-like F2 region (FIG. 27A-B), for example, T-142 to T-145 and T-334 purfied peptides, exhibited detectable levels of antiviral activity, as evidenced 15 by their IC₅₀ values. Further, a number of RSV F1 DP107-like peptides (FIG. 27C-D) exhibited a sizable level of antiviral activity as purified peptides, including, for example, peptides T-124 to T-127, T-131, T-135 and T-137 to T-139, as demonstrated by 20 their low IC_{50} values. In addition, CD analysis FIG. 27A, 27C) reveals that many of the peptides exhibit some detectable level of helical structure.

The results summarized in FIG. 28A-B demonstrate that a number of DP178-like purified peptides exhibit a range of potent anti-viral activity. These peptides include, for example, T-67, T-104, T-105 and T-107 to T-119, as listed in FIG. 28A, and T-665 to T-669 and T-671 to T-673, as listed in FIG. 28B. In addition, some of the DP178-like peptides exhibited some level of helicity.

Thus, the computer assisted searches described, hereinabove, successfully identified viral peptide

domains that represent highly promising anti-RSV antiviral compounds.

18. EXAMPLE: POTENTIAL HUMAN PARAINFLUENZA VIRUS
TYPE 3 DP178/DP107 ANALOGS: CD AND
ANTIVIRAL CHARACTERIZATION

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In the Example presented herein, human parainfluenza virus type 3 (HPIV3) peptides identified by utilizing the computer-assisted search motifs described in the Examples presented in Sections 9 and 15, above, were tested for anti-HPIV3 activity. Additionally, circular dichroism (CD) structural

Additionally, circular dichroism (CD) structural analyses were conducted on the peptides, as discussed below. It is demonstrated that several of the identified peptides exhibit potent antiviral capability. Additionally, it is shown that several of

these peptides exhibit a substantial helical character.

18.1 MATERIALS AND METHODS

Structural analyses: Structural analyses

20 consisted of circular dichroism (CD) studies. The CD
spectra were measured in a 10mM sodium phosphate,
150mM sodium chloride, pH 7.0, buffer at approximately
10mM concentrations, using a 1 cm pathlength cell on a
Jobin/Yvon Autodichrograph Mark V CD

spectrophotometer. Peptide concentrations were determined from A_{280} using Edlehoch's method (1967, Biochemistry <u>6</u>:1948).

Anti-HPIV3 antiviral activity assays: The assay utilized herein tested the ability of the peptides to disrupt the ability of Hep2 cells chronically infected with HPIV3 to fuse and cause syncytial formation on a monolayer of an uninfected line of CV-1W cells. The

more potent the lower the observed level of fusion, the greater the antiviral activity of the peptide.

Uninfected confluent monolayers of CV-1W cells were grown in microtiter wells in 3% EMEM (Eagle Minimum Essential Medium w/o L-glutamine [Bio Whittaker Cat. No. 12-125F], with fetal bovine serum [FBS; which had been heat inactivated for 30 minutes at 56°C; Bio Whittaker Cat. No. 14-501F) supplemented at 3%, antibiotics/antimycotics (Gibco BRL Life Technologies Cat. No. 15040-017) added at 1%, and glutamine added at 1%.

To prepare Hep2 cells for addition to uninfected cells, cultures of chronically infected Hep2 cells were washed with DPBS (Dulbecco's Phosphate Buffered Saline w/o calcium or magnesium; Bio Whittaker Cat.

No. 17-512F) and cell monolayers were removed with Versene (1:5000; Gibco Life Technologies Cat. No. 15040-017). The cells were spun 10 minutes and resuspended in 3% FBS. Cell counts were performed using a hemacytometer. Persistent cells were added to the uninfected CV-1W cells.

The antiviral assay was conducted by, first, removing all media from the wells containing uninfected CV-1W cells, then adding peptides (at the dilutions described below) in 3% EMEM, and 500 chronically HPIV3-infected Hep2 cells per well. Wells were then incubated at 37°C for 24 hours.

On day 2, after cells in control wells were checked for fusion centers, media was removed from the wells, followed by addition, to each well, of approximately $50\mu l$ 0.25% Crystal Violet stain in methanol. Wells were rinsed immediately, to remove excess stain and were then allowed to dry. The number

of syncytia per well were then counted, using a dissecting microscope.

Alternatively, instead of Crystal Violet analysis, cells were assayed with XTT, as described, avove, in Section 17.1.

5 <u>Peptides</u>: The peptides characterized in the study presented herein were:

FIG. 29A); and

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- Peptides 157 to 188, as shown in FIG. 29A, and peptides T-38 to T-40, T-42 to T-46 and T-582, as shown in FIG. 29B. These peptides are derived from the DP107 region of the HPIV3 F1 fusion protein (represented by HPF3 107, as shown in
- Peptides 189 to 210, as shown in FIG. 30A, and T-269, T-626, T-383 and T-577 to T-579, as shown in FIG. 30B. These peptides are primarily derived from the DP178 region of the HPIV3 F1 fusion protein (represented by HPF3 178, as shown in FIG. 30A). Peptide T-626 contains two mutated amino acid resides (represented by a shaded background). Additionally, peptide T-577 represents F1 amino acids 65-100, T-578 represents F1 amino acids 207-242 and T-579 represents F1 amino acids 273-309.

Each peptide was tested at 2-fold serial dilutions ranging from $500\mu g/ml$ to approximately 500ng/ml. For each of the assays, a well containing no peptide was also used.

18.2 RESULTS

The data summarized in FIGS. 29A-C and 30A-B represent antiviral and structural information obtained from peptides derived from the HPIV3 fusion

protein DP107-like region (FIG. 29A-C) and the HPIV3 fusion protein DP178-like region (FIG. 30A-B).

As shown in FIG. 29A-B, a number of the HPIV3 DP107-like peptides exhibited potent levels of antiviral activity. These peptides include, for example, peptides T-40, T-172 to T-175, T-178, T-184 and T-185.

CD analysis reveals that a number of the peptides exhibit detectable to substantial level of helical structure. The CD spectra for one of the peptides,

10 184, which exhibits substantial helicity is summarized in FIG. 29C.

The results summarized in FIG. 30A-B demonstrate that a number of the DP178-like peptides tested exhibit a range of anti-viral activity. These peptides include, for example, peptides 194 to 211, as evidenced by their low IC₅₀ values. In fact, peptides 201 to 205 exhibit IC₅₀ values in the nanogram/ml range. In addition, many of the DP178-like peptides exhibited some level of helicity.

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Thus, the computer assisted searches described,

hereinabove, have successfully identified viral
peptide domains that represent highly promising antiHPIV3 antiviral compounds.

19. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP178/DP107 ANALOGS IN SIMIAN IMMUNODEFICIENCY VIRUS

FIG. 31 represents search results for SIV isolate MM251 (PC/Gene® protein sequence PENV_SIVM2). Both 107x178x4 and ALLMOTI5 search motifs identified two regions with similarities to DP107 and/or DP178.

The peptide regions found by 107x178x4 were located at amino acid residues 156-215 and 277-289.

The peptide regions found by ALLMOTI5 were located at amino acid residues 156-219 and 245-286. Both motifs, therefore, identify similar regions.

Interestingly, the first SIV peptide region (i.e., from amino acid residue 156 to approximately amino acid residue 219) correlates with a DP107 region, while the second region identified (i.e., from approximately amino acid residue 245 to approximately amino acid residue 289) correlates with the DP178 region of HIV. In fact, an alignment of SIV isolate MM251 and HIV isolate BRU, followed by a selection of the best peptide matches for HIV DP107 and DP178, reveals that the best matches are found within the peptide regions identified by the 107x178x4 and ALLMOTI5 search motifs.

region at amino acid residues 242-282 is predicted by the Lupas program. This is similar to the observation in HIV in which the coiled-coil is predicted by the Lupas program to be in the DP178 rather than in the DP107 region. It is possible, therefore, that SIV may be similar to HIV in that it may contain a coiled-coil structure in the DP107 region, despite such a structure being missed by the Lupas algorithm. Likewise, it may be that the region corresponding to a DP178 analog in SIV may exhibit an undefined structure, despite the Lupas program's prediction of a coiled-coil structure.

- 20. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP178/DP107 ANALOGS IN EPSTEIN-BARR VIRUS
- 30 The results presented herein describe the identification of DP178/DP107 analogs within two

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different Epstein-Barr Virus proteins. Epstein-Barr is a human herpes virus which is the causative agent of, for example, infectious mononucleosis (IM), and is also associated with nasopharyngeal carcinomas (NPC), Burkitt's lymphoma and other diseases. The virus 5 predominantly exists in the latent form and is activated by a variety of stimuli.

FIG. 32 depicts the search motif results for the Epstein-Barr Virus (Strain B95-8; PC/Gene® protein sequence PVGLB EBV) glycoprotein gp110 precursor (gp115). The 107x178x4 motif identified two regions 10 of interest, namely the regions covered by amino acid residues 95-122 and 631-658. One PZIP region was identified at amino acid residue 732-752 which is most likely a cytoplasmic region of the protein. The Lupas algorithm predicts a coiled-coil structure for amino acids 657-684. No ALLMOTI5 regions were identified.

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FIG. 33 depicts the search motif results for the Zebra (or EB1) trans-activator protein (BZLF1) of the above-identified Epstein-Barr virus. This protein is a transcription factor which represents the primary mediator of viral reactivation. It is a member of the b-ZIP family of transcription factors and shares significant homology with the basic DNA-binding and dimerization domains of the cellular oncogenes c-fos and C/EBP. The Zebra protein functions as a homodimer. 25

Search results domonstrate that the Zebra protein exhibits a single region which is predicted to be either of DP107 or DP178 similarity, and is found between the known DNA binding and dimerization regions of the protein. Specifically, this region is located at amino acid residues 193-220, as shown in FIG. 33. The Lupas program predicted no coiled-coil regions.

21. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP178/DP107_ANALOGS_IN_MEASLES_VIRUS

FIG. 34 illustrates the motif search results for the fusion protein F1 of measles virus, strain Edmonston (PC Gene® protein sequence PVGLF_MEASE), successfully identifying DP178/DP107 analogs.

The 107x178x4 motif identifies a single region at amino acid residues 228-262. The ALLMOTI5 search motif identifies three regions, including amino acid residues 116-184, 228-269 and 452-500. Three regions containing proline residues followed by a leucine zipper-like sequence were found beginning at proline residues 214, 286 and 451.

The Lupas program identified two regions it predicted had potential for coiled-coil structure, which include amino acid residues 141-172 and 444-483.

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22. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP178/DP107 ANALOGS IN HEPATITIS B VIRUS

FIG. 35 depicts the results of a PZIP motif search conducted on the Hepatitis B virus subtype AYW.

Two regions of interest within the major surface antigen precursor S protein were identified. The first lies just C-terminal to the proposed fusion peptide of the major surface antigen (Hbs) which is found at amino acid residues 174-191. The second region is located at amino acid residues 233-267. The Lupas program predicts no coiled-coil repeat regions.

In order to test the potential anti-HBV antiviral activity of these D178/DP107 analog regions, peptides derived from area around the analog regions are synthesized, as shown in FIG. 52A-B. These peptides represent one amino acid peptide "walks" through the putative DP178/DP107 analog regions. The peptides are

synthesized according to standard Fmoc chemistry on Rinkamide MBHA resins to provide for carboxy terminal blockade (Chang, C.D. and Meinhofer, J., 1978, Int. J. Pept. Protein Res. <u>11</u>:246-249; Fields, G.B. and Noble, R.L., 1990, Int. J. Pept. Protein Res. <u>35</u>:161-214).

Follwing complete synthesis, the peptide aminoterminus is blocked through automated acetylation and the peptide is cleaved with trifluoroacetic acid (TFA) and the appropriate scavengers (King, D.S. et al., 1990, Int. J. Pept. Res. 36:255-266). After cleavage, the peptide is precipitated with ether and dried under vacuum for 24 hours.

The anti-HBV activity of the peptides is tested by utilizing standard assays to determine the test peptide concentration required to cause an acceptable (e.g., 90%) decrease in the amount of viral progeny formed by cells exposed to an HBV viral inoculum.

Candidate antivial peptides are further characterized in model systems such as wood chuck tissue culture and animal sytems, prior to testing on humans.

20 23. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP178/DP107 ANALOGS IN SIMIAN MASON-PFIZER MONKEY VIRUS

The results depicted herein illustrate the results of search motifs conducted on the Simian Mason-Pfizer monkey virus. The motifs reveal DP178/DP107 analogs within the enveloped (TM) protein

GP20, as shown in FIG. 36.

The 107x178x4 motifs identifies a region at amino acid residues 422-470. The ALLMOTIS finds a region at amino acid residues 408-474. The Lupas program predicted a coiled-coil structure a amino acids 424-459.

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24. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP178/DP107 ANALOGS IN BACTERIAL **PROTEINS**

The results presented herein demonstrate the identification of DP178/DP107 analogs corresponding to sequences present in proteins of a variety of bacterial species.

FIG. 37 depicts the search motif results for the Pseudomonas aeruginosa fimbrial protein (Pilin). regions were identified by motifs 107x178x4 and ALLMOTI5. The regions located at amino acid residues 30-67 and 80-144 were identified by the 107x178x4 motif. The regions at amino acid residues 30-68 and 80-125 were identified by the ALLMOTI5.

FIG. 38 depicts the search motif results for the Pseudomonas gonorrhoeae fimbrial protein (Pilin). A 15 single region was identified by both the 107x178x4 and the ALLMOTIS motifs. The region located at amino acid residues 66-97 was identified by the 107x178x4 motif. The region located at amino acid residues 66-125 were identified by the ALLMOTI5 search motif. No coiledcoil regions were predicted by the Lupas program.

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FIG. 39 depicts the search motif results for the Hemophilus Influenza fimbrial protein (Pilin). A single region was identified by both the 107x178x4 and the ALLMOTIS motifs. The region located at amino acid residues 102-129 was identified by the 107x178x4 The region located at amino acid residues 102-148 were identified by the ALLMOTI5 search motif. coiled-coil regions were predicted by the Lupas program.

FIG. 40 depicts the search motif results for the Staphylococcus aureus toxic shock syndrome Hemophilus Influenza fimbrial protein (Pilin). A single region

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was identified by both the 107x178x4 and the ALLMOTI5 The region located at amino acid residues 102-129 was identified by the 107x178x4 motif. region located at amino acid residues 102-148 were identified by the ALLMOTIS search motif. No coiled-5 coil regions were predicted by the Lupas program.

FIG. 41 summarizes the motif search results conducted on the Staphylococcus aureus enterotoxin Type E protein. These results demonstrate the successful identification of DP178/DP107 analogs 10 corresponding to peptide sequences within this protein, as described below.

The ALLMOTI5 motif identified a region at amino acid residues 22-27. The 107x178x4 motif identified two regions, with the first at amino acid residues 26-69 and the second at 88-115. A Pl2LZIPC motif search identified two regions, at amino acid residues 163-181 and 230-250.

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The Lupas program predicted a region with a high propensity for coiling at amino acid residues 25-54. This sequence is completely contained within the first region identified by both ALLMOTI5 and 107x178x4 motifs.

FIG. 42 depicts the search motif results conducted on a second Staphylococcus aureus toxin, enterotoxin A. Two regions were identified by the 25 ALLMOTI5 motif, at amino acid residues 22-70 and amino acid residues 164-205. The 107x178x4 motif found two regions, the first at amino acid residues 26-69 and the second at amino acid residues 165-192. A P23LZIPC motif search revealed a region at amino acid residues 216-250. No coiled-coil regions were predicted by the Lupas program.

FIG. 43 shows the motif search results conducted on the E. coli heat labile enterotoxin A protein, demonstrating that identification of DP178/DP107 analogs corresponding to peptides located within this protein. Two regions were identified by the ALLMOTI5 motif, with the first residing at amino acid residues 55-115, and the second residing at amino acid residues 216-254. The 107x178x4 motif identified a single region at amino acid residues 78-105. No coiled-coil regions were predicted by the Lupas program.

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25. EXAMPLE: COMPUTER-ASSISTED IDENTIFICATION OF DP178/DP107 ANALOGS WITHIN VARIOUS HUMAN PROTEINS

The results presented herein demonstrate the identification of DP178/DP107 analogs corresponding to peptide sequences present within several different human proteins.

FIG. 44 illustrates the search motif results conducted on the human c-fos oncoprotein. The ALLMOTI5 motif identified a single region at amino acid residues 155-193. The 107x178x4 motif identified one region at amino acid residues 162-193. The Lupas program predicted a region at amino acid residues 148-201 to have coiled-coil structure.

FIG. 45 illustrates the search motif results conducted on the human lupus KU autoantigen protein P70. The ALLMOTI5 motif identified a single region at amino acid residues 229-280. The 107x178x4 motif identified one region at amino acid residues 235-292. The Lupas program predicted a region at amino acid residues 232-267 to have coiled-coil structure.

FIG. 46 illustrates the search motif results conducted on the human zinc finger protein 10. The

ALLMOTIS motif identified a single region at amino acid residues 29-81. The 107x178x4 motif identified one region at amino acid residues 29-56. A P23LZIPC motif search found a single region at amino acid residues 420-457. The Lupas program predicted no coiled-coil regions.

26. EXAMPLE: POTENTIAL MEASLES VIRUS DP178/DP107
ANALOGS: CD AND ANTIVIRAL
CHARACTERIZATION

In the Example presented herein, measles (MeV)

virus DP178-like peptides identified by utilizing the computer-assisted search motifs described in the Examples presented in Sections 9 and 21, above, are tested for anti-MeV activity. Additionally, circular dichroism (CD) structural analyses are conducted on the peptides, as discussed below. It is demonstrated that several of the identified peptides exhibit potent antiviral capability. Additionally, it is shown that none of the these peptides exhibit a substantial helical character.

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26.1 MATERIALS AND METHODS

Structural analyses: The CD spectra were measured in a 10mM sodium phosphate, 150mM sodium chloride, pH 7.0, buffer at approximately 10mM concentrations, using a 1 cm pathlength cell on a Jobin/Yvon Autodichrograph Mark V CD spectrophotometer. Peptide concentrations were determined from A₂₈₀ using Edlehoch's method (1967, Biochemistry 6:1948).

Anti-MeV antiviral activity syncytial reduction 30 assay: The assay utilized herein tested the ability of the peptides to disrupt the ability of Vero cells

acutely infected with MeV (i.e., cells which are infected with a multiplicity of infection of 2-3) to fuse and cause syncytial formation on a monolayer of an uninfected line of Vero cells. The more potent the peptide, the lower the observed level of fusion, the 5 greater the antiviral activity of the peptide.

Uninfected confluent monolayers of Vero cells were grown in microtiter wells in 10% FBS EMEM (Eagle Minimum Essential Medium w/o L-glutamine [Bio Whittaker Cat. No. 12-125F], with fetal bovine serum [FBS; which had been heat inactivated for 30 minutes at 56°C; Bio Whittaker Cat. No. 14-501F) supplemented at 10%, antibiotics/antimycotics (Bio Whittaker Cat. No. 17-602E) added at 1%, and glutamine added at 1%.

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To prepare acutely infected Vero cells for addition to the uninfected cells, cultures of acutely infected Vero cells were washed twice with HBSS (Bio Whittaker Cat. No. 10-543F) and cell monolayers were removed with trypsin (Bio Whittaker Cat. No. 17-161E). Once cells detached, media was added, any remaining clumps of cells were dispersed, and hemacytometer cell 20 counts were performed.

· The antiviral assay was conducted by, first, removing all media from the wells containing uninfected Vero cells, then adding peptides (at the dilutions described below) in 10% FBS EMEM, and 50-100 25 acutely MeV-infected Vero cells per well. Wells were then incubated at 37°C for a maximum of 18 hours.

On day 2, after cells in control wells were checked for fusion centers, media was removed from the wells, followed by addition, to each well, of approximately 50µl 0.25% Crystal Violet stain in methanol. Wells were rinsed twice with water immediately, to remove excess stain and were then

allowed to dry. The number of syncytia per well were then counted, using a dissecting microscope.

Anti-MeV antiviral activity plague reduction
assay: The assay utilized herein tested the ability
of the peptides to disrupt the ability of MeV to

infect permissive, uninfected Vero cells, leading to
the infected cells' fusing with uninfected cells to
produce syncytia. The lower the observed level of
syncytial formation, the greater the antiviral
activity of the peptide.

Monolayers of uninfected Vero cells are grown as described above.

The antiviral assay was conducted by, first, removing all media from the wells containing uninfected Vero cells, then adding peptides (at the dilutions described below) in 10% FBS EMEM, and MeV stock virus at a final concentration of 30 plaque forming units (PFU) per well. Wells were then incubated at 37°C for a minimum of 36 hours and a maximum of 48 hours.

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On day 2, after cells in control wells were

checked for fusion centers, media was removed from the
wells, followed by addition, to each well, of
approximately 50µl 0.25% Crystal Violet stain in
methanol. Wells were rinsed twice with water
immediately, to remove excess stain and were then
allowed to dry. The number of syncytia per well were
then counted, using a dissecting microscope.

<u>Peptides</u>: The peptides characterized in the study presented herein were peptides T-252A0 to T-256A0, T-257B1/C1, and T-258B1 to T-265B0, and T-266A0 to T-268A0, as shown in FIG. 47. These peptides represent a walk through the DP178-like region of the MeV fusion protein.

Each peptide was tested at 2-fold serial dilutions ranging from $100\,\mu\text{g/ml}$ to approximately $100\,\text{ng/ml}$. For each of the assays, a well containing no peptide was also used.

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26.2 RESULTS

The data summarized in FIG. 47 represents antiviral and structural information obtained via "peptide walks" through the DP178-like region of the MeV fusion protein.

As shown in FIG. 47, the MeV DP178-like peptides exhibited a range of antiviral activity as crude peptides. Several of these peptides were chosen for purification and further antiviral characterization. The IC₅₀ values for such peptides were determined, as shown in FIG. 47, and ranged from 1.35μg/ml (T-257B1/C1) to 0.072μg/ml (T-265B1). None of the DP178-like peptides showed, by CD analysis, a detectable level of helicity.

Thus, the computer assisted searches described, hereinabove, as in for example, the Example presented in Section 9, for example, successfully identified viral peptide domains that represent highly promising anti-MeV antiviral compounds.

27. EXAMPLE: POTENTIAL SIV DP178/DP107 ANALOGS: ANTIVIRAL CHARACTERIZATION

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In the Example presented herein, simian immunodeficiency virus (SIV) DP178-like peptides identified by utilizing the computer-assisted search motifs described in the Examples presented in Sections 9, 12 and 19, above, were tested for anti-SIV activity. It is demonstrated that several of the

identified peptides exhibit potent antiviral capability.

27.1 MATERIALS AND METHODS

Anti-SIV antiviral assays: The assay utilized

5 herein were as reported in Langolis et al. (Langolis,
A.J. et al., 1991, AIDS Research and Human
Retroviruses 7:713-720).

Peptides: The peptides characterized in the
study presented herein were peptides T-391 to T-400,
as shown in FIG. 48. These peptides represent a walk
through the DP178-like region of the SIV TM protein.

Each peptide was tested at 2-fold serial dilutions ranging from $100\mu g/ml$ to approximately 100ng/ml. For each of the assays, a well containing no peptide was also used.

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27.2 RESULTS

The data summarized in FIG. 48 represents antiviral information obtained via "peptide walks" through the DP178-like region of the SIV TM protein.

As shown in FIG. 48, peptides T-391 to T-400 were tested and exhibited a potent antiviral activity as crude peptides.

Thus, the computer assisted searches described, hereinabove, as in for example, the Example presented in Section 9, for example, successfully identified viral peptide domains that represent highly promising anti-SIV antiviral compounds.

28. EXAMPLE: ANTI-VIRAL ACTIVITY OF DP107 AND DP-178 PEPTIDE TRUNCATIONS AND MUTATIONS

The Example presented in this Section represents a study of the antiviral activity of DP107 and DP178

truncations and mutations. It is demonstrated that several of these DP107 and DP178 modified peptides exhibit substantial antiviral activity.

28.1 MATERIALS AND METHODS

- Anti-HIV assays: The antiviral assays performed were as those described, above, in Section 6.1.

 Assays utilized HIV-1/IIIb and/or HIV-2 NIHZ isolates.

 Purified peptides were used, unless otherwise noted in FIGS. 49A-C.
- 10 <u>Peptides</u>: The peptides characterized in the study presented herein were:
 - 1) FIGS. 49A-C present peptides derived from the region around and containing the DP178 region of the HIV-1 BRU isolate.
- Specifically, this region spanned from gp41
 amino acid residue 615 to amino acid residue
 717. The peptides listed contain
 truncations of this region and/or mutations
 which vary from the DP178 sequence amino
 acid sequence. Further, certain of the
 peptides have had amino- and/or carboxyterminal groups either added or removed,
 as indicated in the figures; and
 - 2) FIG. 50. presents peptides which represent truncations of DP107 and/or the gp41 region surrounding the DP107 amino acid sequence of HIV-1 BRU isolate. Certain of the peptides are unblocked or biotinylated, as indicated in the figure.

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Blocked peptides contained an acyl N-terminus and an amido C-terminus.

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activity.

28.2 RESULTS

Anti-HIV antiviral data was obtained with the group 1 DP178-derived peptides listed in FIG. 49A-C. The full-length, non-mutant DP178 peptide (referred to in FIG. 49A-C as T20) results shown are for 4ng/ml.

In FIG. 49A, a number of the DP178 truncations exhibited a high level of antiviral activity, as evidenced by their low IC50 values. These include, for example, test peptides T-50, T-624, T-636 to T-641, T-645 to T-650, T-652 to T-654 and T-656. T-50 10 represents a test peptide which contains a point mutation, as indicated by the residue's shaded background. The HIV-1-derived test peptides exhibited a distinct strain-specific antiviral activity, in that none of the peptides tested on the HIV-2 NIHZ isolate demonstrated appreciable antti-HIV-2 antiviral

Among the peptides listed in FIG. 49B, are test peptides representing the amino (T-4) and carboxy (T-3) terminal halves of DP178 were tested. The amino terminal peptide was not active (IC₅₀>400 µg/ml) whereas 20 the carboxy terminal peptide showed potent antiviral activity (IC₅₀= $3\mu g/ml$). A number of additional test peptides also exhibited a high level of antiviral activity. These included, for example, T-61/T-102, T-217 to T-221, T-235, T-381, T-677, T-377, T-590, T-25 378, T-591, T-271 to T-272, T-611, T-222 to T-223 and T-60/T-224. Certain of the antiviral peptides contain point mutations and/or amino acid residue additions

In FIG. 49C, point mutations and/or amino and/or carboxy-terminal modifications are introduced into the DP178 amino acid sequence itself. As shown in the

which vary from the DP178 amino acid sequence.

figure, the majority of the test peptides listed exhibit potent antiviral activity.

Truncations of the DP107 peptide (referred to in IG. 50 as T21) were also produced and tested, as shown in FIG. 50. FIG. 50 also presents data concerning blocked and unblocked peptides which contain additional amino acid residues from the gp41 region in which the DP107 sequence resides. Most of these peptides showed antiviral activity, as evidenced by their low IC₅₀ values.

Thus, the results presented in this Section demonstrate that not only do the full length DP107 and DP178 peptides exhibit potent antiviral activity, but truncations and/or mutant versions of these peptides can also possess substantial antiviral character.

29: EXAMPLE: POTENTIAL EPSTEIN-BARR DP178/DP107
ANALOGS: ANTIVIRAL CHARACTERIZATION

In the Example presented herein, peptides derived from the Epstein-Barr (EBV) DP-178/DP107 analog region of the Zebra protein identified, above, in the Example presented in Section 20 are described and tested for anti-EBV activity. It is demonstrated that among these peptides are ones which exhibit potential antiviral activity.

29.1 MATERIALS AND METHODS

Electrophoretic Mobility Shift Assays (EMSA):
Briefly, an EBV Zebra protein was synthesized
utilizing SP6 RNA polymerase in vitro transcription
and wheat germ in vitro translation systems (Promega
Corporation recommendations; Butler, E.T. and
Chamberlain, M.J., 1984, J. Biol. Chem. 257:5772;
Pelham, H.R.B. and Jackson, R.J., 1976, Eur. J.

Biochem. 67:247). The in vitro translated Zebra protein was then preincubated with increasing amounts of peptide up to 250 ng/ml prior to the addition of 10,000 to 20,000 c.p.m. of a 32P-labeled Zebra response element DNA fragment. After a 20 minute incubation in the presence of the response element, the reaction was analyzed on a 4% non-denaturing polyacrylamide gel, followed by autoradiography, utilizing standard gelshift procedures. The ability of a test peptide to prevent Zebra homodimer DNA binding was assayed by the peptide's ability to abolish the response element gel migration retardation characteristic of a protein-bound nucleic acid molecule.

Peptides: The peptides characterized in this study represent peptide walks through the region containing, and flanked on both sides by, the DP178/DP107 analog region identified in the Example presented in Section 20, above, and shown as shown in FIG. 33. Specifically, the peptide walks covered the region from amino acid residue 173 to amino acid residue 246 of the EBV Zebra protein.

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Each of the tested peptides were analyzed at a range of concentrations, with 150ng/ml being the lowest concentration at which any of the peptides exerted an inhibitory effect.

29.2 RESULTS

The EBV Zebra protein transcription factor contains a DP178/DP107 analog region, as demonstrated in the Example presented, above, in Section 20. This protein appears to be the primary factor responsible for the reactivation capability of the virus. A method by which the DNA-binding function of the Zebra virus may be abolished may, therefore, represent an

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effective antiviral technique. In order to identify potential anti-EBV DP178/DP107 peptides, therefore, peptides derived from the region identified in Section 20, above, were tested for their ability to inhibit Zebra protein DNA binding.

The test peptides' ability to inhibit Zebra protein DNA binding was assayed via the EMSA assays described, above, in Section 28.1. The data summarized in FIG. 51A-B presents the results of EMSA assays of the listed EBV test peptides. 10 peptides represent one amino acid "walks" through the region containing, and flanked on both sides by, the DP178/DP107 analog region identified in the Example presented in Section 20, above, and shown as shown in FIG. 33. As shown in FIG. 51A-B, the region from which these peptides are derived lies from EBV Zebra protein amino acid residue 173 to 246. A number of the test peptides which were assayed exhibited an ability to inhibit Zebra protein homodimer DNA binding, including 439, 441, 444 and 445.

Those peptides which exhibit an ability to 20 inhibit Zebra protein DNA binding represent potential anti-EBV antiviral compounds whose ability to inhibit EBV infection can be further characterized.

> EXAMPLE: IDENTIFICATION OF RSV DP107/DP178 ANALOGS WITH REDUCED BINDING AFFINITY

In the example presented herein, peptides derived from the RSV DP178 analog T112 are described and tested for binding affinity to the DP107-like domain of the RSV F1-protein. Particular peptides are 30 identified that have a reduced binding affinity for their DP107-like target, and key amino acid residues

are identified the confer high binding affinity to the native peptide (i.e., to T112). Such peptides are useful, e.g., in screening assays such as those described above in Section 5.6.1 to identify compounds which inhibit or disrupt the interaction between DP107 and DP178, and in providing guidance for generation of additional peptides exhibiting reduced affinity binding.

30.1 MATERIALS AND METHODS

10 A maltose binding fusion protein of the RSV F1protein (MF5.1) was constructed using methods similar
to those described in Section 8.1.2, supra, for
construction of the M41 fusion protein. Specifically,
the DNA sequence corresponding amino acid residues
142-302 of the RSV F1 protein was amplified by PCR and
cloned into the Xmn I site of the expression vector
pMal-p2 (New England Biolab) to give MF5.1. These
amino acid residues correspond to the extracellular
domain of the RSV F1 protein including its DP107
region but excluding the DP178 region.

The peptides characterized in the study presented herein were: T122, T800, T801, T802, T803, T804, T805, T806, T807, T808, T809, T810, T811, T1669, T1670, T1671, T1672, T1673, T1680, T1681, T1682, T1683 and T1684, as shown in FIG. 53. T112 represents the DP178-like region of the RSV F1 protein. The other peptides characterized are modified DP178 proteins derived from T112.

Cell fusion assays were performed with each of the peptides as described in Section 17 above. The binding affinity of each peptide was also measured in a competitive binding assay described in Section 5.6.1

above, wherein the concentration of each peptide necessary to bind to the M5.1 fusion protein (i.e., the B_{50} value), and thereby disrupt binding of biotin labeled T112 (T888) to the fusion protein, was measured.

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30.2 RESULTS

T112 is a 35 amino acid residue peptide that corresponds to amino acid residues 482-516 of the RSV F1 protein and has the following amino acid sequence:

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VFPSDEFDASISQVNEKINQSLAFIRKSDELLHNV

The peptide represents the DP178-like region of the RSV F1 protein and has substantial antiviral activity against RSV as discussed in Section 17.2 above and shown in FIG. 28A.

T112 analogs were generated according to at least three different strategies to generate peptides based on T112 that would still bind to the DP107-like domain of the RSV F1 protein but with a lower binding affinity. First, a truncated peptide was generated, reducing the length of the peptide from 35 to 28 amino acid residues. Specifically, the truncated peptide, which is referred to herein as T67, had the amino acid sequence:

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DEFDASISQVNEKINQSLAFIRKSDELL

corresponding to amino acid residues 486-213 of the F1 fusion protein. The binding affinity of the peptide to the DP107-like domain of F1 protein was determined according to the methods described in Section 5.6.1

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> above. The truncated peptide had a binding affinity (5 nM) that was five times lower than that of the full length T112 peptide (2 nM).

As part of a second strategy, the peptides identified as T800 through T811 in FIG. 53 were 5 synthesized to identify particular amino acids in T112 that contribute to a larger part of that peptide's binding affinity. As a whole, these alanine substitutions represent an "alanine-scanning" type walk across the sequence of T112.

Each of the peptides synthesized had a change of 10 three consecutive amino acid residues in the T112 sequence to three alanine residues. Each peptide was tested for its ability to inhibit the binding of the native peptide (i.e., of T112) in a competitive binding assay as described in Section 5.6.1 above. 15 The results are also shown in FIG. 53. In particular, the peptides T802, T804, T807 and T810 had significantly reduced affinity for the DP107-like target, suggesting that the regions containing amino acid residues 488-490, 494-496, 503-505 and 512-514 of the RSV F1 protein (amino acid residues 7-9, 13-15, 22-24 and 31-33, respectively, of T112), contribute significantly to the high binding affinity of T112 for its DP107-like target in the RSV F1 protein.

The peptides T1669-T1673 and T1680 through T1684 25 were then synthesized, each of which contains a single alanine substitution at one of the above-listed amino acid residue positions of T112. The binding affinity of these peptides for their DP107-like target can also be determined by means of the same routine screening assays, thereby identifying individual amino acid residues which affect binding affinity of T112.

In addition, an additional novel peptide, referred to as T786, was generated by modifying various amino acid residues in the T112 sequence which were identified, using standard principles of protein and design, as affecting properties such as binding affinity, solubility and biological stability. Specifically the following amino acid residue substitutions were made: F₂ - Y, S₂₁ - A, F₂₄ - Y and S₂₈ - A, wherein the subscript numerals indicate the amino acid residue position in T112. The resultant peptide, which is referred to herein as T786, thus had the amino acid sequence:

VYPSDEFDASISQVNEKINQALAYIRKADELLHNV

The binding affinity of this novel peptide for the DP107 target was found to be 19 nM, i.e., approximately ten-fold less than the binding affinity of T112.

The data demonstrates that peptides having a reduced binding affinity for a DP107 target (i.e., for an HR1 domain) may be readily found by modifying a DP178 peptide such as T112, e.g., by means of the routine techniques and assays described herein. Further, the techniques and assays identify key amino acid residues which may be used to construct and identify other reduced affinity peptides.

31. EXAMPLE: IDENTIFICATION OF HIV DP107/DP178
ANALOGS WITH REDUCED BINDING
AFFINITY

In the example presented herein, peptides derived 30 from DP178, which is also referred to as T20, are described and tested for binding affinity to the DP107

domain of the HIV gp41. Particular peptides are identified that have a reduced binding affinity for their DP107 target, and key amino acid residues are identified the confer high binding affinity to the native peptide (i.e., to T20). Such peptides are useful, e.g., in screening assays such as those described above in Section 5.6.1 to identify compounds which inhibit or disrupt the interaction between DP107 and DP178.

Specifically, the peptides identified as T813 and 10 T868 through T878 in FIG. 53 were synthesized to identify particular amino acids in T20 (DP178) that contribute to a greater part of that peptide's binding affinity. Each of the peptides synthesized had a change of three consecutive amino acid residues in the T20 sequence to three alanine residues. The antiviral 15 activity of each peptide was assayed in cell fusion assays as described in Section 6.1.3, above. binding affinities of the peptides were also measured in a competitive binding assay described in Section 5.6.1 above, wherein each peptides ability to disrupt the binding of either biotin (T83) or fluorescein (T1342) labeled DP178 (T20) to the M41A178 fusion protein described in Section 8, above, was measured. The binding affinity of each peptide to the peptide referred to as T764

25 (GSTMGARSMTLTVQARQLLSGIVQQNNLLRAIEAQQH) also measured using circular dichroism to monitor the amount of secondary structure (i.e., helicity) adopted by the peptides. T764 is a peptide which represents the DP107 target domain of DP178 (T20).

The results are provided in FIG. 54. In particular, the peptides T813, T878, T874-T876 and

T871 have significantly reduced affinity for the DP107 region, suggesting the regions corresponding to the substituted amino acid residues in those peptides contribute significantly to the high binding affinity of T20. The peptides T1627-T1632, T1650-T1653 and 5 T1656-T1665 were then synthesized. Each of these peptides contains a single alanine substitution at one of the amino acid residues in one of the regions identified as contributing significantly to the high binding affinity of T20. Identical assays which measured the binding affinity of these peptides identified four essential residues (I $_{646}$, Q $_{652}$, Q $_{653}$ and N_{656} , with the subscript numerals indicating the residue position in the HIV-1LAI gp41 amino acid sequence) in which alanine-substitution completely prevented binding to the DP107 domain, as well as five 15 residues (L_{641} , I_{642} , I_{645} , E_{657} and L_{663} , with the subscript numerals indicating the residue) in which alanine-substitution position in the HIV-1 ap gp41 amino acid sequence) that reduced the binding affinity but did not actually block binding to the DP107 20 domain.

The data demonstrates that peptides having a reduced binding affinity for a DP107 target (i.e., for an HR1 domain) may be readily found by modifying a DP178 peptide such as T20, e.g., by means of the routine techniques and assays described herein. Further, the techniques and assays identify key amino acid residues which may be used to construct and identify other reduced affinity peptides.

The present invention is not to be limited in scope by the specific embodiments described which are intended as single illustrations of individual aspects of the invention, and functionally equivalent methods and components are within the scope of the invention.

Indeed, various modifications of the invention, in addition to those shown and described herein will become apparent to those skilled in the art from the foregoing description and accompanying drawings. Such modifications are intended to fall within the scope of the appended claims.

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WHAT IS CLAIMED IS:

A method for identifying a compound that inhibits the formation of or disrupts a DP107/DP178
 complex comprising:

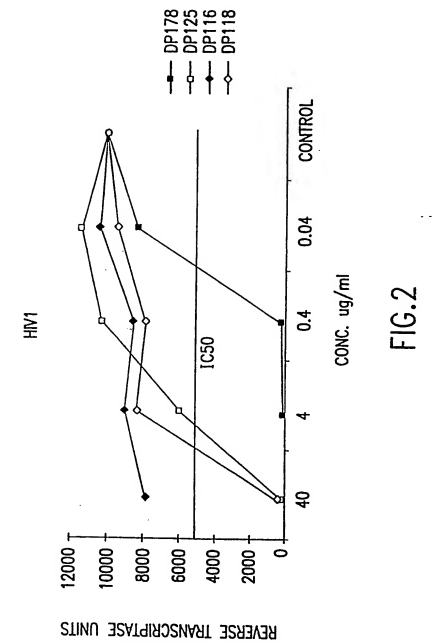
- (a) preparing, both in the presence and in the absence of a test compound, a reaction mixture containing a DP107 peptide and a DP178 peptide under conditions and for a time sufficient to allow formation of a DP107/DP178 complex; and
- (b) detecting the formation of a DP107/DP178 complex both in the presence and in the absence of the test compound,
- wherein the formation of a DP107/DP178 complex in the absence, but not in the presence of the test compound indicates that the compound inhibits the formation of or disrupts a DP107/DP178 complex.
- 2. The method of Claim 1, wherein the DP107 peptide or the DP178 peptide is a modified DP107 or DP178 peptide.
 - 3. The method of Claim 2 wherein the modified DP107 or DP178 peptide has a reduced binding affinity.

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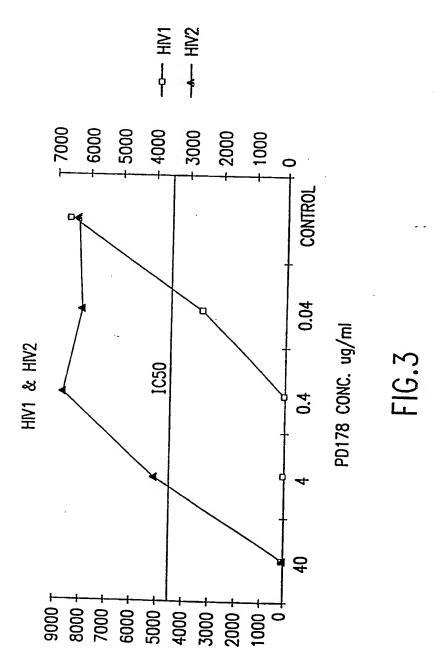
LQARILAVERYL	DP116 (SEQ ID:9)
CGGNNLLRAIEAQQHLLQLTVMG1KQLQAR1LAVERYL	DP125 (SEQ ID:8)
QQLLDVVKRQQEMLRLTVWGTKNLQARVTAIEKYLKDQ	DP118 (SEQ ID:10)
SSESFTLLEGWNNWKLQLAEGWLEQINEKHYLEDIS	DP180 (SEQ ID:2)
LEANISQSLEQAQIQQEKNAMELQKLNSMDVFTNML	HIV2NIHZ (SEQ ID:7)
LEANISKSLEQAQIQQEKNAYELQKLNSWDIFGNWF	HIV2ROD (SEQ ID:6)
YTSL IYSLLEKSQTQQEKNEQELLELDKWASLWNWF	HIV1MN (SEQ ID:5)
YTG I IYNLLEESONQOEKNEOELLELDKWANLWNWF	HIV1RF (SEQ ID:4)
YTNTIYNLLEESONQOEKNEGELLELDKWASLWNWF	HIV1SF2 (DP-185; SEQ ID:3)
HIV1LAI (DP-178; SEQ ID:1) YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNF	HIVILAI (DP-178; SEQ ID:1)

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Number of Syncytia/well: concentration in μg/m1 (micrograms/ml)										
DP178	10	5	11	0.2	0.1	0.05	0.025	0.0125	Control	
Syncylia		_	_				_			
HIVILAI	0	0	0	0	0	0	0	0	67	
HIVIM	0	0	0	0	0	ND	ND	ND	34	
HIVIRF	0	0	0	0	0	ND	ND	ND	65	
HIV1SF2	0	0	0	0	0	ND	ND	ND	58	
DP125	10	5	1	0.2	0.1	0.05	0.025	0.0125	Control	
Syncylia				<u> </u>	0.1	0.00	0.020	0.0120	00:121 01	
HIVILAL	0	0	54	69	80	75	79	82	67	
HIVIMN	Ö	Ö	30	36	ND	ND	ND	ND	34	
HIVIRF	Ō	Ö	67	63	ND	ND	ND	ND	65	
HIV1SF2	Ō	0	9	66	ND	ND	ND	ND	58	
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•										
DP116	10	5	1	0.2	0.1	0.05	0.025	0.0125	Control	
Syncytia										
HIVILAT	75	ND	ND	ND	ND	ND	ND	ND	67	
HIVIMN -	35	ND	ND	ND	MD	ND	ND	ND	34	
HIV1RF	81	ND	ND	ND	ND	ND	ND	ND	65	
HIV1SF2	81	ND	ND	ND	ND	МD	MD	ND	58	

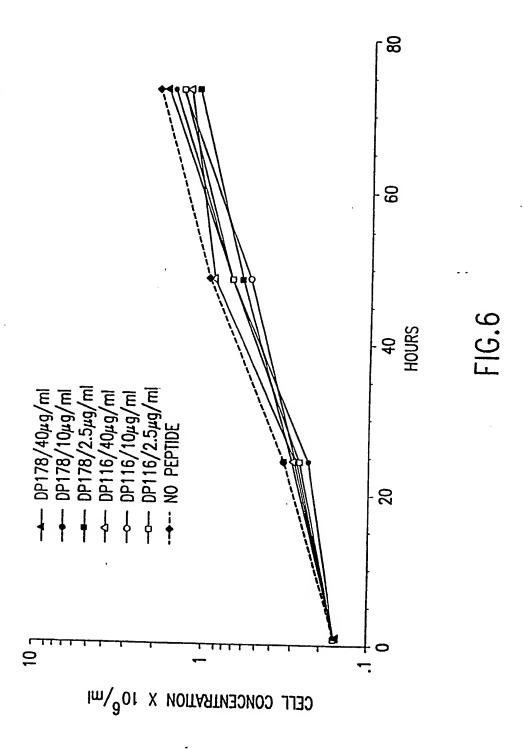
FIG.4A

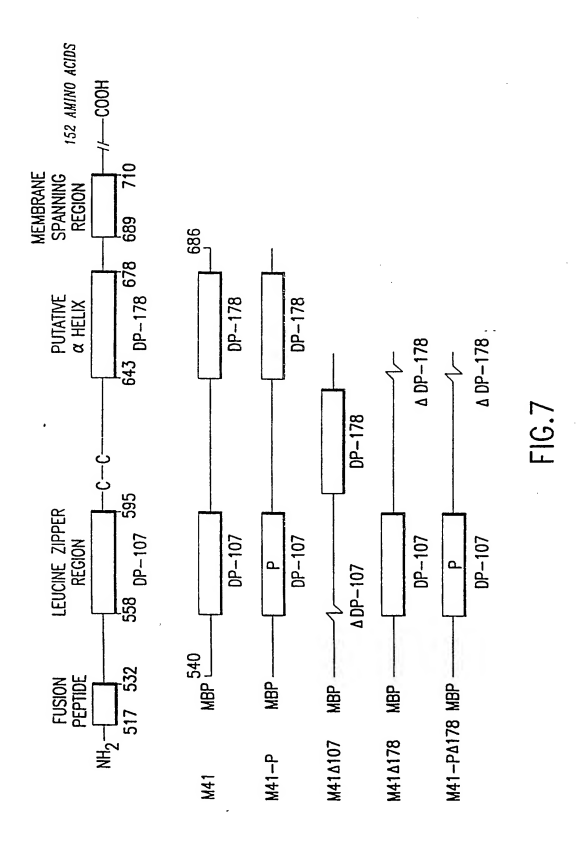
<u>DP180</u>	40	20	10	5	2.5	1.25	0.625	0.3125	Control
Syncylia HIV1LAI	50	>45	>45	>45	>45	>45	>45	>45	58
DP185	40	20	10	5	2.5	1.25	0.625	0.3125	Control
Syncylia HIVILAI	0	0.	0	0	0	0	0	ND	60

FIG.4B

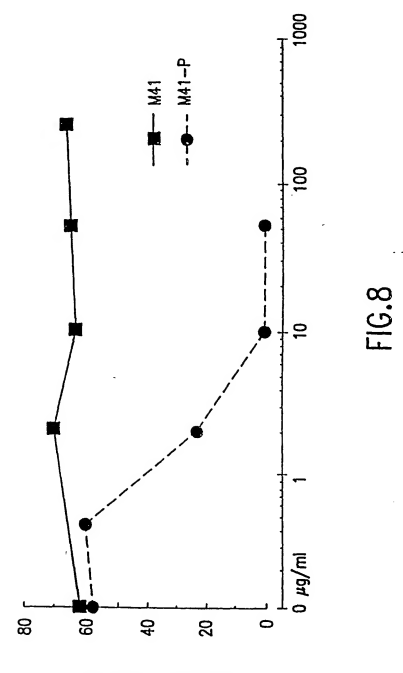
HIV1											
	Number	of	Syncyl	ia/well:	conce	ntration	in ng/ml	(nanograms/ml)			
DP178	20	10	5	2.5	1.25	0.625	0.3125	Control			
Syncytio HIVI	0	0	0	0	0	14	20	48			
DP116	20	10	5	2.5	1.25	0.625	0.3125	Control			
Syncylia HIVI	ND	48	ND	ND	ND	ND	ND	ND			
	ні V2										
	Number	of	Syncyt	io/well:	conce	ntralion	in μg/ml	(micrograms/ml)			
DP178	20	10	5	2.5	1.25	0.625	0.3125	Control			
Syncylia HIV2	50	54	55	57	63	77	78	76			
DP116	20	10	5	2.5	1.25	0.625	0.3125	Control			
Syncylia HIV2	ND	58	ND	ND	ND	ND	ND	ND			

FIG.5

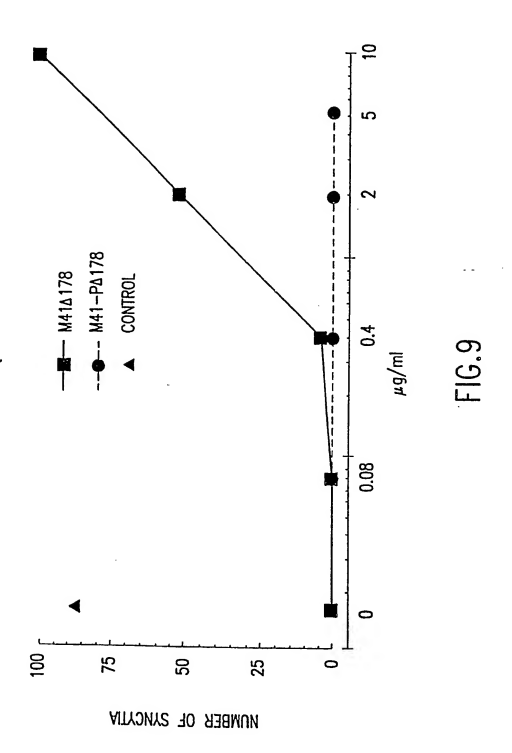




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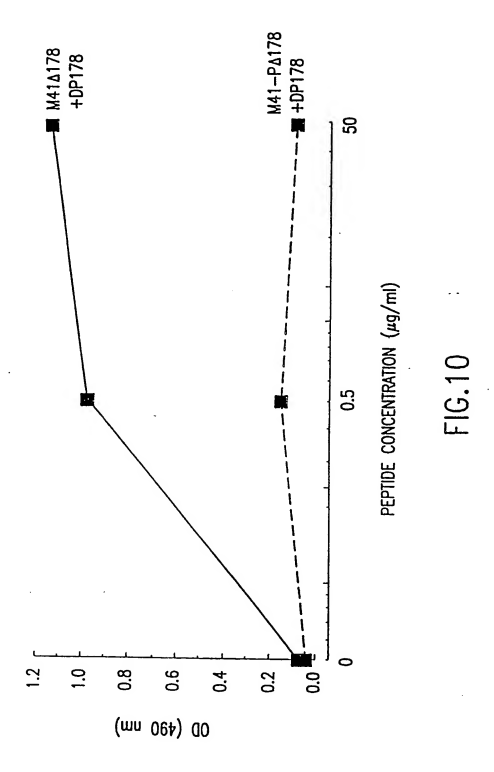


NUMBER OF SYNCYTIA

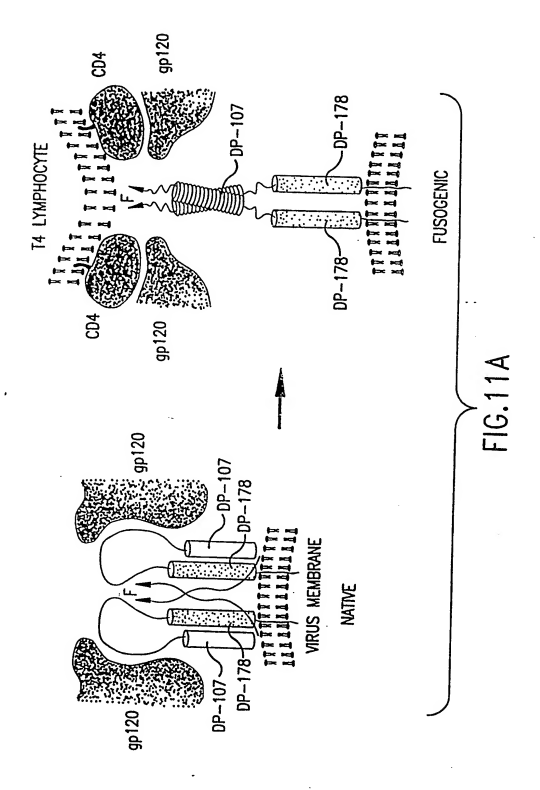


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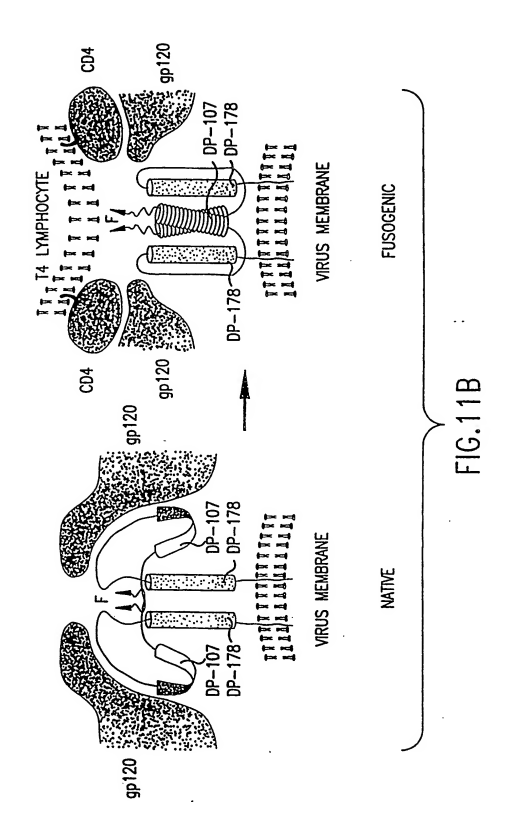


FIG. 12

						Positic	ions					Mobile
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GCN4 · (gcn4 yeast)	MK 0	1 E 0	K V E E	LLSK	Η N	LENE	VAR	L K K	L		1	[LANV] (CEGIAPIW)
C-FOS (fos_human)	101	L 0 A	E	0 I E D E	K S A	10 1 5	EIAN	L K F		 		[KI I] {CECHINDRAWY}
C-JUN (tap1_human)	I A R	IARLEE	K V K	VKTLKAO	N S	LAS1	ANM	2		 		[AII NV] {CDECHII DVW?
C-MYC (myo_human)	E 0 X	L 1 S	EEOL	LEKR	R E O	LKHK	CLEOLR	02				[FIR] {ACECADAWY}
FLU LOOP 36	IEKTN	TNE	K F H O	I E K E F S E V E G R I O D L	FI SE	VEGR	0 0 1	w				[FILTV] {ACFLIPPTW#}

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Motifs		[1LQT] {CF IMPSTY}	[1LOTV] {COF IMPST}	[1LQTV] {CDF1MPST}	[EKLNOV] {CDFKAPSVY}	[EKLNOV] {CFIOAPS}	[EKLNQV] {CFKNPS}	[EKLQY] {ACFGNPRVMY}	[EKLOWY] {CFGMPRVY}	[EFKLOHY] {CFGNPRVY}	[E11 NOSY] {ACFGNPRVMY}	[ETLNOSHY] (CFGNPRVY)	[EFILNOSIM] (CFGIRPRY)
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	Sednence	DP-107 (env_hv1bru)L1=0	DP-107 (env_hv1bru)L1=D	DP-107 (env_hv1bru)L1=D	OP-107 (env_hv1bru)L2=D	DP-107 (env_hv1bru)L2=D	OP-107 (env_hv1bru)L2=D	DP-178 (env_hv1bru)Y1=A	2	8	œ	DP-178 (env_hv1bru)Y1=0	20
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Hybrid Motif		[[LMOTV] {CF IMPT}	[ILMADIV] (CFIMPT)	[ILMNOTV] {CFINPT}	[EKLMNOV] {CFNP}	[EKLIMOV] (CFINP)	[EKLMOV] (CFMP)
Parent Motif	[LMNV] {CFGIMPTW}	[ILQT] {CF IMPSTY}	[1LQTV] (CDF IMPST)	[ILOTV] {COFINPST}	[EKLNOV] {CDFKAPSVY}	[EKLNOV] (CFKAPS)	[EKLNOV] {CFKNPS}
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Sequence	GCN4 (gen4 yeast)	0P-107 (env_hv1bru)L1=	OP-107 (env_hv1bru)∟1=	0P-107 (env_hv1bru)l.1€	DP-107 (env_hv1bru)L2=0	DP-107 (env_hv1bru)L2=0	DP-107 (env_hv1bru)L2=0 N N L L R A 1 E
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FIG. 14

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Hybrid Notif			[EKLIMOYY] {CFGIAPH}	[EKLINDWIY] (CFGIP)	[EFKLIANDYNY] {CFCAP}	[EILNOSY] {ACFGAPRVAY} [EILMNOSYY] {GFGAPA}	[ETHANDSWAY] (CFGAP)	I HI SICII E EISIO NIOLO E KINIE OLEIL L'EILID KINIA S L'ININ NIFI [EFILMOSNY] (CFONPRYY) [EFILMADSWY] (CFONP)
Parent Molif		[LMNY] {CFGIMPTW}	[EKLQY] {ACFGMPRWMY}	[EKLOHY] (CFGHPRVY)	[EFKLOMY] {CFCLAPRYY}	[EILNOSY] {ACFGAPRYM	[EILNOSHY] (CFCMPRVY	[EFILNDSWY] (CFCHPRVY
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	equence	GCN4 (gcn4 yeast)	P-178 (env_hv1bru)Y1=	P-178 (env_hv1bru)Y1=/	DP-178 (env_hv1bru)Y1=A Y	OP-178 (env_hv1bru)Y1=D	P-178 (env_hv1bru)Y1=	DP-178 (env_hv1bru)Y1=0
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Hybrid Holif	[FEIKLINGSTVMY] {GWP}	
Porent Motif	[ILQTV] {CDF IMPST} [EKLNOV] {CPKAPS} [EFKLOMY] {CFGAPRVY} [EFLLNOSMY] {CFGAPRVY}	[FILTV] {ACFUAPTWII}
[0]	AQQHLLQLTVWGIKQLQARIILAVERYLKDQ QQHLLQLTVWGIKQLQARILAVERYLKDQ EESQNQQEKNEQELLELDKWASLWWWF SLIEESQNQQEKNEQELLELDKWASLWWF	
IS DI A	V W G I K O L O A R I L A V E R Y L W G I K O L O A R I L A V E R Y L K N E O E L L E L D K W A S L W N W F O E K N E O E L L E L D K W A S L W	1 0 0 L E K Y
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Sequence	0P-107 (env_hv1bru)L1=D NNLLRAITE 0P-107 (env_hv1bru)L2=D NNLLRAITE DP-178 (env_hv1bru)Y1=A YTSLIHSLIH 0P-178 (env_hv1bru)Y1=D YTSLIH	FLU LOOP 36

	1	,		i	Posilions	ons					Parent Motif	Hybrid Motif
Sequence	A 0	~	0	0 Y	0	A	0	₹	٥			
GCN4 (gen4 yeast)	MK OLL	E D K V	1 5 6 1 1	SKNY	H E N	EVAR	L K K				[LIMNY] {CFG[IMPTW]	
0P-107 (env_hv1bru)[1=0	NN	L R A	E A 0 0	H [[0]	1 / 1	clik o	LOAS	8 1 1 A	V F R Y	O X		
DP-178 (env_hv1bru)Y1=A	YTSLIMSLIE	I M S	I E E S	ONOO E	<u> </u>	OFLU	10 1	K W A S	¥ N ★	EESONOOEKNEOELLELDKWASLWNWF	[EFKLOHY] [CFGHPRYY]	[EF IKLIANDTWAY] {CFIAP}
										_		
GCN4 (gen4 yeast)	MX OLE	₹ 0 ×	<u> </u>	SKINY	N H	EVAR	ואאו				[LMNV] {CFGINPTW}	
OP-107 (env_hv1bru)L1=0	NN	RA	E A 0 0	10 1 1 H	1 V W (3 K 0	LOAE	RIIL A	VERY	l K D O	[1LOTY] (COF IMPST)	
0P-178 (env_hv1bru)Y1=0	Y T S L II H S	S [H S I	EESON	100E	C NE O) X	WASL	SLIFE ESONO OF KINE OF LLELD KINA SLIMN WIF	[EF ILNOSIY] (CFCNPRVY)	[EF][MORSTWY] {CFILP}
		_										
GCN4 (gan4 yeast)	MK OLE	0 K	<u> </u>	SKNYH	I E N E	VAR	LXXL				[LIAN] (CFGIAPTH)	
DP-107 (env_hv1bru)L2=0	NNLLRAIEAO	A IE	A OOH 1	O O H L LOL TIVING IKO LOAR ILAVERYIKDO	S <u>**</u> ∧	K O L	OAR I	I A V	FRY	0 U	[EKLNOV] (CFIO.PS)	
OP-178 (env_hv1bru)Y1=A	YI SI	H S	I EES	3 Nolo E	X X	E	FIDK	V X	3 2	, ,	[FFKI CHY] [CFCLIPRY)	[FFKI LANDWRY] {CFLIP}
				· ·		, , ,	; , ,		: : :			[
	MK OLE	. 0 K	П П	S K N Y H	<u> </u>	V A R	K K K				[LIAN] (CFG IMPTH)	
	NNLLR	V V	A OOH L	1011	[5 <u>₩</u> 6	K O	OAR I	Y	K 4 1	- X	[EKLNOV] {CFINIPS}	
	YTSLIHSI	112	H SIC II	ILIEESONOOEKNEOELLELDKWASIWAN	OOEX	NEO	- L	2	I S V	7 X	· [EF IL NOSHY] {CFCLIPRY}	[FF KI LANSWAY] (PS.LP)
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FIG. 17

Hybrid Molif										[AEF IKLIANORSTM	{\ruser_{\curuer_{\ruser_{\ruser_{\curuer_{\ruser_{\curuer_{\curuer_{\curuer_{\curuer_{\curuer_{\curuer_{\ruser_{\curuer_{
Parent Motif		[LMN] {CFCINPTH}	[1LQTV] (COF IMPST)	[EKLNOV] (CFNAPS)	[EFKLOHÝ] (CFCHPRW)	[EFILNOSIM] {CFCHPRVM}	[IKLT] (CFCHIMPRYMY)	[AILNY] (COFGHILPWRY)	[ELR] {ACFGNPYNY}	[FILTY] {ACFLIAPTIVITY}	
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	Sequence	GCN4 (gen4 yeast)	OP-107 (env_hv1bru)L1:	DP-107 (env_hv1bru)L2=	DP-178 (env_hv1bru)Y1=	DP-178 (env_hv1bru)Y1=D Y T S	C-FOS (fas_human)	C-JUN (top1_human)	C-MYC (myo_human)	FLU 100P 36	

FIG. 18

WO 01/51673 PCT/US00/35727

Fusion

YALLMOTI5₩

Peptide

4107x178x44

Y.....FLGFLG A AGSTMGARSM TLTVQARQ ALLSGIVOOO DP107-NNL

LRAIEAOOIIL LOLTVIYGIKO LOARILAYER YLKDO-DP107 QLLG** I WGC

4<u>107x178x4</u>4

YALLMOTISY

LVS Coiled-Coil

SGKLICT TAVP *WNASWS NKSLEQIWNN MTWM *E *WDREINN DP178-

YTSLIHSL IEESONOOEK NEOELLELDK* WASLIVNIVE-DP178 NI

+ Transmembrane Region +

TNWLWYIK + IF IMIYGGLYGLRIVEAVLSIY NRVRQGYS + PL

+P23LZIPC+

SFQTHLPTPR GPDR #PEGIEE EGGERDRDRS IRLVNGSLAL IWDDLRSL# CL

YALLMOTISY

4107x178x44

F VSYHRLRDLL LIVTRIVELL GRRGW *EALKY WWNLLOYWSO

ELKNSAVSLL NAT ↑ AIAVAEG TDRVIEVVQG A♥ CRAIRHIPR

RURQGLERIL L

Peptide

4107x178x44

V.....DGEL LGVGSAIAS GVA AYSKYLILLEGEYNKIKSA

+P1&12LZIPC+

LLSTNKAYYS LSNGYSYLTS KYLDLKNYID KQ * LL +PIVNKQ

4107x178x44

SC *SISNIETY I+ EFOOKNNRLLEITREFSYNAG* VITI'VSTMLINSELLSL

P1&12LZIPC

YALLMOTI5Y

INDM →PI →TNDQ KKLMSNNVQI V→ RQQSYSI→ MS IIKEEVŁAYV

VQ▼ LPLYGVID TPCWKLIITSP LCTTNTKEGS NICLTRTDRG WYCDNAGSVS

FFPQAETCKV QSNRVFCDTM NSLTLPSEIN LCNVDIFNPK

YDCKIMTSKT DVSSSVITSL GAIVSCYGKT KCTASNKNRG

IIKTFSNGCDYVSNKGMDTV SVGNTLYYVN KQEGKSLYVK G

+P7, 12, & 23LZIPC+

4107x178x44

YALLMOTISY

EPHNFYDPLVF +PSDE +FDASISOYNEKINOSLAF *I+ RKSDELL+

+ Transmembrane Region +

LINVNA + GK STTN + IMITTI LIVILVILLS LIAYGLLLY + C+

KARSTPYTLS KDOLSGINNI AFSN

Fusion

Peptide VALLMOTI5 * * 107x178x4 *

.....FLGFLG YAAGTA MGAAA ATALTYOSQULLAGUQQQKNLLAAY

+107x178x4+

EAO+ QQM +LKLTIWGYKNLNARYTALEKYLEDOARLN+ AWGY CA

LYS Coilcd-Coil

YALLMOTI5Y +107x178x4+

WKQVCHTIVP WQWNNRTPDW VNNMT *WLE *WERQISYLEGNIT

4107x178x44

TOLEEARAOEEKNLD * AYOKLSS* WSDFWSW* FDF *SKWLN +ILK

+ Transmembrane Region +

IGFLDYLGHGLRLLYTY + YS + CIARVRQGYS PLSPQHIIH WKGQPDNAEG

PGEGGDKRKN SSEPWQKESG TAEWKSNWCK RLTNWCSISS IWLYNS

YALLMOTI5Y

▼CLTL LVHLRSAFQY IQYGLGELKA AAQEAVVALA RLAQNAGYQIWL▼

ACRSAYRA IINSPRRVRQ GLEGILN

WO 01/51673

Fusion #107x178x4*

1'eptide VALLMOTI5♥ *LVS Coiled-Coil*

.....FAG *VYL AGVALGVATA AQITAGIALHQ **SNLNAQAIQ

SLRTSLEOSNKAIEEIREATOETYIA* YOGYODY* VNNEL* VP

YALLMOTI5 Y

4107x178x44

4P6 & 12LZIPC4

AMQHMSCELVGQRLGLRLLRYYTELLSIFGPSLRD +PISA +VEISIOALIYAL

GGEILIKILEKLGYSGSD → MIAILESRGIKTKI → THVDLPGKF IILŞISY

+P1 & 12LZIPC+

+PTILSEVKGVIVHRLEAV+ SYNIGSQEWYTTVPRYIATNGYLISNFDESSCVFVS

ESAICSQNSL YPMSPLLQQE IRGDTSSCAR TLVSGTMGNK FILSKGNIVA

NCASILCKCY STSTIINQSP DKLLTFIASD TCPLVEIDGA TIQVGGRQYP

LVS Coilcd-Coil

YALLMOTISY

+P12 & 23LZIPC+

DMVYEGKVAL G *PAISLD *RL*DYGTNLGNALKKLDDAKYLJ*

+ Transmembrane Region +

DSS+ NOILETYRRS+ SFN + FGSLL SYPILSCIAL ALLLLIYCC+

K RRYQQTLKQH TKVDPAFKPD LTGTSKSYVR SL

Fusion YALLMOTISY

Peptide ★107x178x4★

▼......FIGAI IGSVALGVA TAAQITAASA LIQANQNAAN ★ILRLKESITA

TIEAVIIEVTDGLSOLAYA + VG KM + QQFVNDQFNNTAQELDCIKITQQV

♥ ALLMOTI5 ♥
GVELNLYLTELTTV FGPQITSPAL ▼TQLTIQALYNAGGNMDYLLTKLGVG

+P1 & 12LZIPC+
NNQLSSLIGSGLIT GN▼ +PILYDSQT QLLGIQVTLP SVGNLNNMRATYLET

LSVST TKGFASALVP KVVTQVGSVI EELDTSYCIE TDLDLYCTRI VTFPMSPGIY
.
SCLNGNTSAC MYSKTEGALT TPYMTLKGSV IANCKMTTCR CADPPGIISQ

LYS Coiled-Coil

NLDISTELGNY NNSISNALDK LEESNSKLDK YNYKLTSTSA +LIT YIA

membrane Region +
LTAISLVCGILSLV + LACYLMY + KQKAQQKTLLWLGNNTLGQMRATIKM

Fusion

YALLMOTIS#

Peptide

107x178x4 *LVS Coilcd-Coil*

.....ITGGY

*IG *TIALG *YATSAQITAAYALYEAKQARSDIEKLKE

AIRDTNKAYOSVOSSIGNLIYAIKSVO* DYVNKE** IVPSIARLGCEAAG

YALLMOTI5Y

4107x178x44

LOLGIALTOH *YSELTNIFGDNIGSLOEKGIKLOGIASLYRTNITEY*

+P5 & 12LZIPC+

IFTTSTVDKYDIYDLLFTESIKVRVIDVDLNDYSITLQVRL +PLLTRLLNTQIYR

VDSISYNI+ QNREWYI+ PLPSHIMTKGAFLGGADVKECIEAFSSYIC

PSDPGFVLNHEMESCLSGNISQCPRTVVKSDIVPRYAFVNGGVVANCITT

TCTCNGIGNRINQPPDQGVKIITHKECNTIGINGMLFNTNKEGTLAFYTP

YALLMOTISY

4107x178x44

+P6 & 23LZIPC+

NDITLNNSVALD +PIDI +SIELN +KAKSDLEESKEWI+ RRSNOKL+

+ Transmembrane Region +

DSIGNWHOSSTT + HIV + LIM HILFHNYTH+ HAVKYY + R

IQKKNIKVDQN DKPYVLTNK

Fusion
Peptide
......GLEGAL AGFIENGWEGMIDGWYGFRHQNSEGTG

4107x178x44

YALLMOTI5Y

LYS Coilcd-Coil

*Q *AADLKST *QAAIDQINGKLNRYIEKTNEKFHQIEKEFSEYEGRIQ

DLEKYYEDTKIDL* YSYNAELLYALENQITI* DLT* DSEMNKLFEKTR
RQLRENAEEMGNGCFKIYHKCDNACIESIRNGTYDHDVYRDEALNNRFQIKG
VELKSGYKDWILWISFAISCFLLCVVLLGFIMWACQRGNIRCNICI

PCT/US00/35727

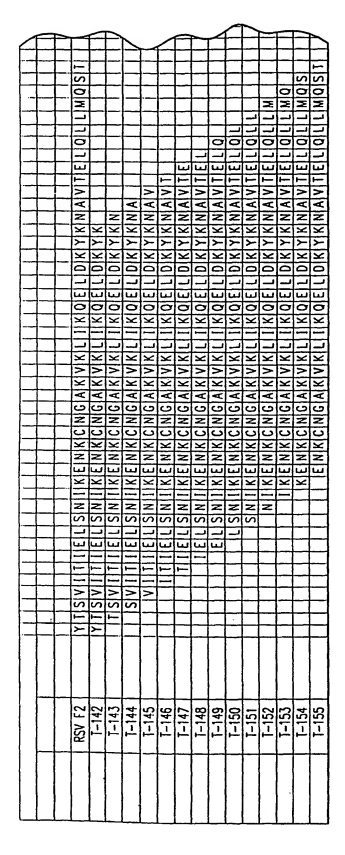


FIG.2/A

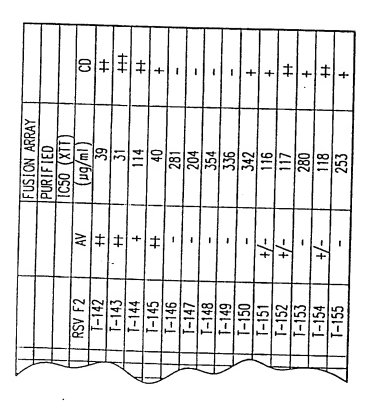


FIG.27B

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1 EL SIN 1 K EN K C N G T D A K V K L 1 K O E L D K Y		>500
T-24 EINKCNGTDAKIVKLIKQELDKYKNAVTEL		>500
I I I I I I I DAKVKL		>500
O N		>500
T-27		>500
T-68 VISIKIGIYISIAILIRITIGIWIYITISIVIIITII IEILISINIIIKIEINI IIIIIIIIIIIIIIIIIIIIIIIIIIII		165
T-334 A F 1 R K S D E L L H N V		76
YTISIVITTIELISINITKENKUNGTDAKVKL		>500
TSVITIELSNIIKENKUNG TDAKVKL		NOT TESTED
SVITIELSINIKENKUNGTOAKVKLIKOE		>200
N S		>500
T-375	[[M 0 S	>500
T-575 A V S K G Y L S A L R T G W Y T S V ! T E L S N T K E N K U N G T D A		>100

FIG.2/C

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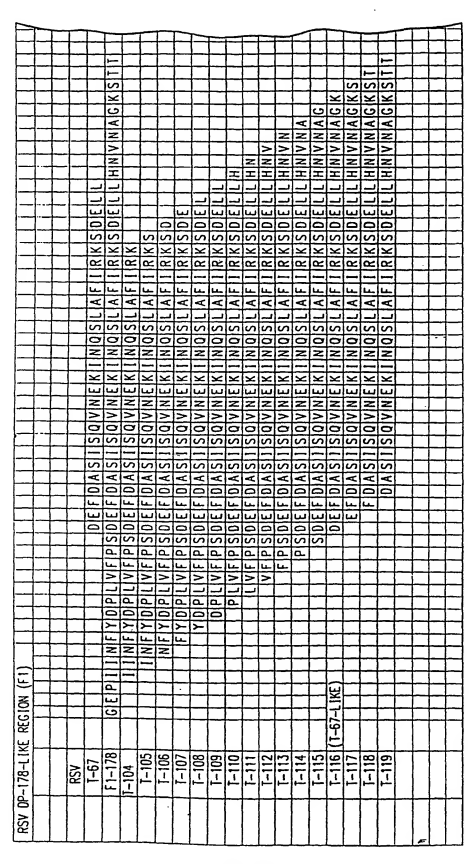
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1-70 V N K T K S A L L S T N K A V V S L S N G V S V L T S K	349
T-66 NIDIQIK KILIMISIN IN V 0 1 V R O 0 1 W S 1 W S 1 1 K E E 1 1 1	005<
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FIG. 27F



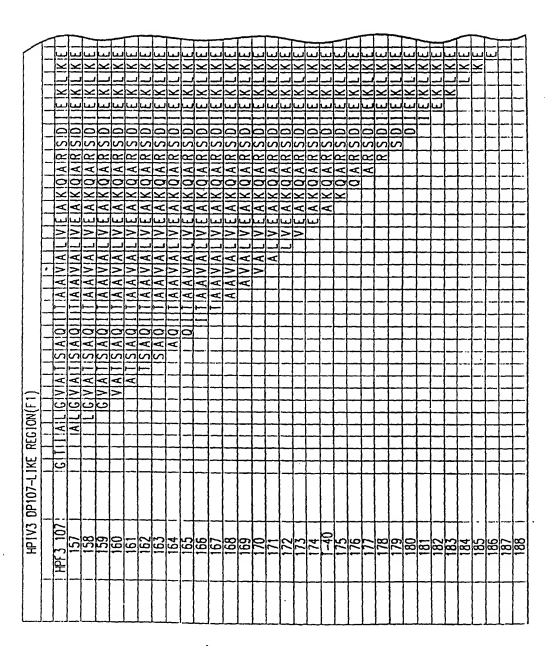
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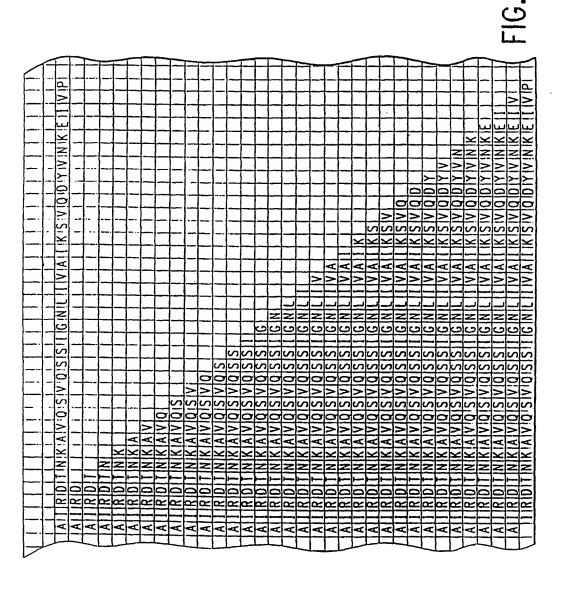
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			RSV	1-67	F1-178	1-104	T-105	T-106	/ T-107	1-108	1-109	1-110	1-111	1-112	T-113	T-114	1-115	T-116	1-117	T-118	T-119		

FIG. 28B

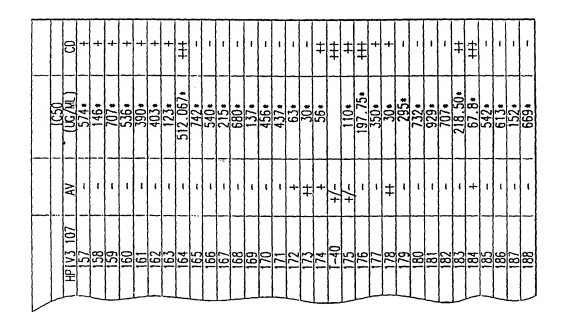
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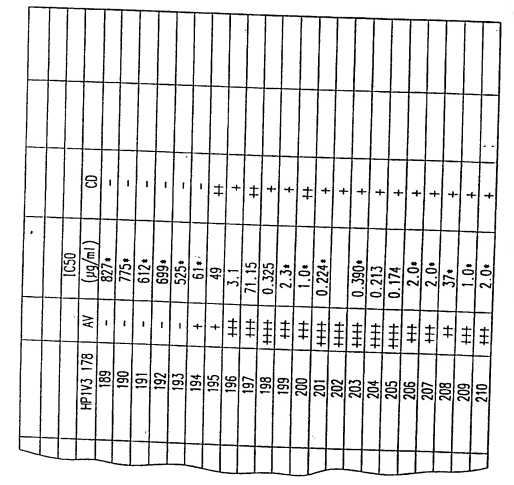
FIG. 29D

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T-269 TRUNCATED 201				I E W	EWIRRISINIOKLIDISII	KLOSII		457.500UC/ML
T-626 205 MUTANT	11011811	11 10 11 15 11 1E 1C 1N 1K 1A 1K 1S 10 1C 1E 1S 1K 1E 1W 11 1K 1K 1S N 10 1K 1C 10 1S 1 1 1G N W H	SIBITOS	K E W I	KKSNO	KILIDISITIG	H M N	209.589NG/ML
T-383 RIMIK OLIEIDIKIVIEIELLIS	뒬	EWIRRSNOKLDSI	SII					NOT DONE
T-577 0 0 0 0 1 K 0 Y K R L L 0 R L	IIPLYDGL	PLYDGLRQKDVIVSNQESN	IVSNOE	S				133.793UG/ML*
1 579 YE FE T M 1 FC D M 1 C S	-	L L L L L L L L L L	0	- - - - -				107 1771F ARE
200	<u> </u>	2 - 2				 		101.1700/
T-579 TISTITLOVRLPLLTR	블	NTIQIIYRVDISIISYNIIONREWY	YNIONR	E W Y				NOT DONE

FIG. 30C

FUSION PEPTIDE

♥ALLMOTI5**♥**

±107x178x4**±**

....RNKRGVFVLGFLGFLATAGSAMGAAS AY XXXXAOSRTLLAGIVOOQOO

LLDVVKRQQELLRLTVWGTKNLQTRVTAIEKYLKDQAQL4NAWG♥ CAF

▼ALLMOTIS▼

*LVS PREDICTED COILED-COIL

RQVCHTTVPWPNASLTPDW *NND ▼TWQEWERKVDFLEENITALLEEAQIQQ

<u>\$107x178x4</u>\$
EKNMY <u>\$ELQKLNSWD</u>* <u>VF</u>♥ <u>GNXXXXXXXXXXXXXXXXXXXXXXXX</u>\$

. IYIVMLAKLRQGYRPVFSSPPSYFQXTHTQQDPALPTREGKEGDGGEGGGNSSWP WQIEYIHF

MTRRRVI.SVVVLLAALACRLGAQTPEQPAPPATTVQPTATRQQTSFPFRVCELSSHGDLFRFSSD

10CPSFGTRENHTEGLLMVFKONIIPYSF ★ KVRSYTKIVTNILIYNGWYADSVTNRHE ★
EKFSVDSY ETDQMDTIYQ CYNAVKMTKD GLTRVYVDRD GVNITVNLKP TGGLANGVRR

YASQTELYDA PGWLIWTYRT RTTVNCLITD MMAKSNSPFD FFVTTTGQTV EMSPFYDGKN

KETFHERADS FHVRTNYKIV DYDNRGTNPQ GERRAFLDKG TYTLSWKLEN RTAYCPLQHW

QTFDSTIATE TGKSIHFVTD EGTSSFVTNT TVGIELPDAF KCIEEQVNKT HEKYEAVQD

RYTKGQEAIT YFITSGGLLL AWLPLTPRSL ATVKNLTELT TPTSSPPSSP SPPAPSAARG

STPAAVLRRR RRDAGNATTP VPPTAPGKSL GTLNNPATVQ IQFAYDSLRR QINRMLGDLA

RAWCLEQKRQ NMVLRELTKI NPTTVMSSIY GKAVAAKRLG DVISVSQCVP VNQATVTLRK

<u>♦107x178x4</u> EIHVYNDYHH FKTIELDGIA TLQTFISLNT <u>♦SLIENIDFASLELYSRDEQRASNVFD</u> *LE♠

SMRVPGSETM CYSRPLVSFS FINDTKTYEG QLGTDNEIFL TKKMTEVCQA TSQYYFQSGN

LVS PREDICTED COILED COIL TM Potential
GIFREYNFQAQNIAGLRKDLDNAVSN* GRNQ FVDGLGELMDSLGSVG OSITN

♣P12LZIPC♣

TM Potential
LVSTVGGLFSSLVSGFISF FK N &PFGGMLILVLVAGVVILVISL TRRTRQMS

QQPVQMLYPG IDELAQQHAS GEGPGINPIS KTELQAIMLA LHEQNQEQKR AAQRAAGPSV

ASRALQAARDRFPGLRRRRY HDPETAAALL GFAETEF

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MMDPNSTSLD VKFTPDPYQV PFVQAFDQAT RVYQDLGGPS QAPLPCVLWP VLPEPLPQGQ LTAYHVSTAP TGSWFSAPQP APENAYQAYA APQLFPVSDI TQNQQTNQAG GEAPQPGDNS TVQTAAAVVF ACPGANQGQQ LADIGVPQPA PVAAPARRTR KPQQPESLEE CDSELEI

@DNA_BINDING@

<u> 107x178X4</u>

+DIMERIZATION+

OKRY KNRVASRKCRAK

♠FK@ Q

+LLOHYREVAAAKSSENDRLRLLLKQ♠

MCPSLDVD+ SI IPRTPDVLHE DLLNF

FUSION

PEPTIDE

♥ALLMOTI5♥

LVS_COILED-COIL

FAG

♥VVLAGAALGVATAAQITAGIALHQSML*NSOAIDNLRASLETTN

<u>OAIEAIRGAGOEMI</u>*LAVQGVQDYINN♥ ELIPSMNQLSCDLIGQKLGLKLLRYYT

♣P23LZIPC♣

♣P6,12LZIPC♣

★107X178X4★

♥ALLMOTI5♥

LIFSLEGPSLRD ♣PISA ♠♥EISIQLSYALGGDINKV♣ LEKLGYSGGDL♣

♣P1.12LZIPC♣

LGILES♠ RGIKARI♥ THVDTESYFIVLSIAY ♣PTLSEIKGVIVHRLEGV♣ SY

NIGSQEWYTTVPKYVATQGYLISNFDESSCTFMPEGTVCSQNALYPMSPLLQECL

RGSTKSCARTLVSGSFGNRFILSQGNLIANCASILCKCYTTGTIINQDPDKILTYIAA

♣P23LZIPC♣

♣P12LZIPC♣

♥ALLMOT15♥

LVS_COILED-COIL

DHCPVVEVNGVTIQVGSRRYPDAVYLHRIDLGP ♣P ♥IS *LERLDVGTNLGN

♦TRANSMEMBRANE REGION

AJAKLEDAKELL♣ ESSDOI*L♣ RSMK ♦GLSSTSIVYILI♥ AVCLGGLIGIP

<u>ALICCC</u>♦ RGRCNKKGEQVGMSRPGLKPDLTGTSKSYVRSL

Pre S1 and Pre S2
MGQNLSTSNPLGFFPDHQLDPAFRANTANPDWDFNPNKDTWPDANKVGAGAFG
LGFTPPHGGLLGWSPQAQGILQTLPANPPPASTNRQSGRQPTPLSPPLRNTHPQAM
QWNSTFFHQTLQDPRVRGLYFPAGGSSSGTVNPVLTTASPLSSIFSRIGDPALN

MAJOR SURFACE ANTIGEN(HBs)
FUSION
PEPTIDE
#P12 & 23LZIPC#

MENITSG FLG *PLL VI.QAGFFLLTRILTI PQSLDSWWTSLNFLGGTTVCLG

♣P12 & 23LZIPC♣
QNSQSPTSNHSPTSCPPTC ♣PGYRWMCLRRFIIFLFILLLCLIFLLVLLDYQGML♣
PVCPLIPGSSTTSTGPCRTCMTTAQGTSMYPSCCCTKPSDGNCTCIPIPSSWAFGKF

◆TRANSMEMBRANE REGION◆
LWEWASARFSWLS ◆LLVPFVOWFYGLSPTVWLSVI◆ WMMWYWGPSI.

- **♦**TRANSMEMBRANE REGION**♦**
- **+YSILSPFLPLLPIFFCLWVYI+**

WO 01/51673 PCT/US00/35727

FUSION ▼ ALLMOTI5 ♥ <u>\$107x 178x4</u>

PEPTIDE *LVS COILED COIL

AIQLIPLFVG LGI ▼TTAVSTGAAGLGVS <u>\$1T</u> *<u>QYTKLSHOLISDV</u>

QAISSTIQDLQDQVDSLAEVVLQ* NRRGLDLLTAE♠ QGGI♥

CLALQEKCCFYANKSGIVRDKIKNLQDDLERRRRQLIDNPFWTSFHG

FLPYVMPLLGPLLCLLLVLSFGPIIFNKLMTFIKHQIESIQAKPIQVHYII

TRANSMEMBRANE REGION RLEQEDSGGSYLTLT......????????????????????????

MKAQKGFTLI ELMIVVAIIG ILAAIAPGQ

±107x178x4±

YALLMOTI5Y

♦♥<u>YODYTARTQVTRAVSEVSALKTAAESAILEGKEIYSSA</u>**•** T♥

PK DTQYDIGFT

±107x178x4**±**

YALLMOTI5Y

≜♥<u>ESTLLDGSGKSQIQVTDNQDGTVELVATLGKSSGS</u>**±** AIKGAVITSR♥

KNDGV WNCKITKTPT AWKPNYAPAN CPKS

MNTLQKGFTL IELMIVIAIV GILAAVALPA YQDYTARAQV

SEAILLAEGQ KSAVTEYYLN HGIWP

- ±107x178x4±
- YALLMOTI5Y
- **★Y**KDNTSAGVASSSSIKGKYVKEVKVENGVVTAT

MNSSNVNKEIQGKKLSLWAKRQDGSVKW♥

FCGQP VTRNAKDDTV TADATGNDGK IDTKHLPSTC RDNFDAS

MKKTLLGSLI LLAFAGNVQA DINTETSGKV TFFGKVVENT

CKVKTEHKNL SVVLNDVGKN SLSTKVNTAM PTPFTITLQN

CDPTTANGTA NKANKVGLYF Y

- **±**107x178x4**±**
- **♥**ALLMOTI5♥
- **♦♥**SWKNYDKENNFTLKNEQTTADYATNYNI

QLMESNGTKAISVVGKETE**▼**

DIF MHTNNNGVAL NQTIIPNNAHI SGSTQLTTGT NELPLHFIAQ

YYATNKATAG KVQSSVDFQI AYE

MNKKLLMNFF IVSPLLLATT ATDFTPVP

- ±107x178x4±
- **♥**ALLMOT15♥
- **♦VLSSNOIIKTAKASTNDNIKDLLDWYSSGSDTFTNS♦V**

EVLDNSL GSMRIKNTDG SISLIIFPSP YYSPAFTKGE KV

- **±**107x178x4**±**
- **♦**DLNTKRTKKSQIITSEGTYIHFQISGVT**♦**

N TEKLPTPIEL PLKVKVHGKD SPLKYG

- ♣P12LZIPC♣
- **♣PKFDKKQLAISTLDFEIRHQLTQI♣**

HGLYRSSDKT GGYWKITMND GSTYQSDLSK KFEYNTEKPP

INIDEIKTIE AEIN

♥ALLMOTI5♥

MKKTAFILLL FIALTLTTSP L ▼VNG

±107x178x4±

- *LVS PREDICTED COILED-COIL*
- *S <u>♦ EKSEEINEKDLRKKSELQRNALSNLRQIY</u>* <u>YYNEKAITENKESDD</u>

QFLENTLL♥ FKG FFTGIIPW

- ±107x178x4±
- **<u>♦YNDLLVDLGSKDATNKYKGKKVDLYGAY</u>**

YGYQCAGGTPNKTACMYGGVTLIIDN NRLTEEKKVP INLWIDGKQTTV

- ・ ◆P12LZIPC◆
 - ♣PIDKVKTSKKEVTVQELDL♣ QARHYLHGK FGLYNSDSFGGKVQ

♣P12LZIPC♣

RGLIVF HSSEGSTVSY DLFDAQGQY ♣P DTLLRIYRDN KTINSENLHI♣

DLYLYTT

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¥ALLMOTIS¥ MKKTAFTLLL FIALTLTTSP L ¥VNGS

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- **♦EKSEEINEKDLRKKSELOGTALGNLKOIYYYNEKAKTENKESHD♦ Q♥**

FLQIITILFKG FFTDIISWYND LLVDFDSKDI VDKYKGKKVDLYGAYY

GYQC AGGTPNKTAC MYGGVTLIIDN NRLTEEKKVPINL WLDGKQNTV

♣P12LZIPC♣

♣P ♥L ♠ETVKTNKKNVTVOELDLOARRYL♣ OEKYNLYN♠

SDVFDGKVQR♥ GLIVF HTSTE

♣P23LZIPC♣

♣PSVNYDLFGAQGQYSNTLLRIYRDNKTINSENMHI**♣** DIYLYTS

MKNITFIFFILLASPLYANGDRLYRADSRPPDEIKRFRSLMPRGNEYFDRGT

- **♥**∧LLMOTI5**♥**
- **♥**QMNINLYDHARGTQTGFVRYDDGYV
- ±107x178x4±
- **♦STSLSLRSAHLAGOYILSGYSLTIYIVI♦** ANMFNVNDVISVY♥

SP HPYEQEVSAL GGIPYSQIYG WYRVNFGVID ERLHRNREYR

DRYYRNLNIA PAEDGYRLAG FPPDHQAWRE EPWIHHAPQG

CGDSSRTITG DTCNE

- **♥**ALLMOTI5♥
- **▼**ETQNLSTIYLREYQSKVKRQIFSDYQSEVDIYNRIRDEL**▼**

MMFSGFNADY EASSSRCSSA SPAGDSLSYY HSPADSFSSM GSPVNAQDFC TDLAVSSANF IPTVTAISTS PDLQWLVQPA LVSSVAPSQT RAPHPFGVPA PSAGAYSRAG VYKTMTGGRA

LVS PREDICTED COILED-COIL
QSIGRRGKVE QLSPEEEEKR RIRRE *RNKMA AAK

±107x178x4**±**

- **♥**∧LLMOTI5♥
- ♥CRNRRREL <u>♠TDTLOAETDOLEDEKSALOTEIANLLKEKEKL</u>♥

EFILAAHR* PACKIPDDL GFPEEMSVAS LDLTGGLPEV
ATPESEEAFT LPLLNDPEPK PSVEPVKSIS SMELKTEPFD
DFLFPASSRP SGSETARSVP DMDLSGSFYA LPLLNDPEPK
PSVEPVKSIS SMELKTEPFD DFLFPASSRP SGSETARSVP
DMDLSGSFYA GSSSNEPSSD SLSSPTLLAL

SGWESYYKTEGDEEAEEQEENLEASGDYKYSGRDSLIFLVDASKA MFESQSEDELTPFDMSIQCIQSVYISKIISSDRDLLAVVFYGTEKDKNS VNFKNIYVLQELDNPGAKRILELDQFKGQQGQKRFQDMMGHGSDY SLSEVLWVCANLFSDVQFKMSHKRIMLFTNEDNPHGNDSAKASRAR TKAGDLRDTGIFLDLMHLKKPGGFDISLFYRDIISIAEDED

±107x178x4**±**

♥ALLMOTI5♥

LVS PREDICTED COILED-COIL

¥LRVH *FEE \$SSKLEDLLRKVRAKETRKRALSRLKLKLNKDIV* ISV

GIYNLVQKAL♥ KPPPIKLYRETN♠ EPVKTKTRTFNTSTGGLLLPSDTKR

SQIYGSRQIILEKEETEELKRFDDPGLMLMGFKPLVLLKKHHLRPSLFVYPE ESLVIGS STLFSALLIKCLEKEVAALCRYTPRRNIPPYFVALVPQEEELDDQK IQVTPPGFQLVFLPFADDKRKMPFTEKIMATPEQVGKMKAIVEKLRFTYRS DSFENPVLQQIIFRNLEALALDLME

♣PI2LZIPC♣

♣PEQAVDLTLPKVEAMNKRL**♣** GSLVDEFKELVYPPDYNPEGKVTKR KHDNEGSGSKRPKVEYSEEELKTHISKGTLGKFTVPMLKEACRAYGLKSG LKKQELLEALTKHFQD

GGGALSPQIISAVTQGSIIKNKEGMDAKS

±107x178x4±

♥ALLMOTI5♥

▼ LTAWSRTLYTFKDVFYDFTREEWKLLDT AQQIVYRNVMLENYKNLVSLGYQLT

KPDVILRLEKGEEPWLVEREIHQETHPD
SETAFEIKSSVSSRSIFKDKQSCDIKMEGMARNDLWYLSLEEVWKCR
DQLDKYQENPERIILRHQLIHTGEKPYECKECGKSFSRSSHLIGHQKT
HTGEEPYECKECGKSFSWFSHLVTHQRTHTGDKLYTCNQCGKSFVH
SSRLIRHQRTHTGHKPYECPECGKSFRQSTHLILHQRTHVRVRPYECN
ECGKSYSQRSHLVVHHRIHTGLKPFECKDCGKCFSRSSHLYSHQRTH
TGEKPYECIIDCGKSFSQSSALIVHQRIHTGEKPYECCQCGKAFIRKN
DLIKIIQRIIIVGAETYKCNQCGIIFSQNS

♣P23LZIPC**♣**

♣PFIVHQIAHTGEQFLTCGNQCGTALVNTSNLIGQTNHI**♣** RENAY

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FIG. 47B

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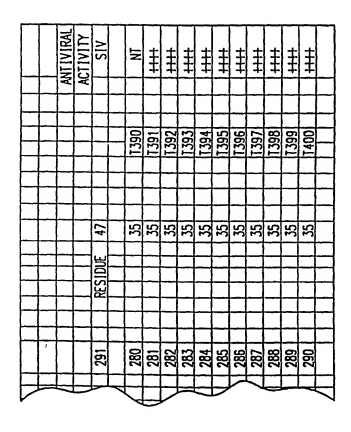


FIG.48B

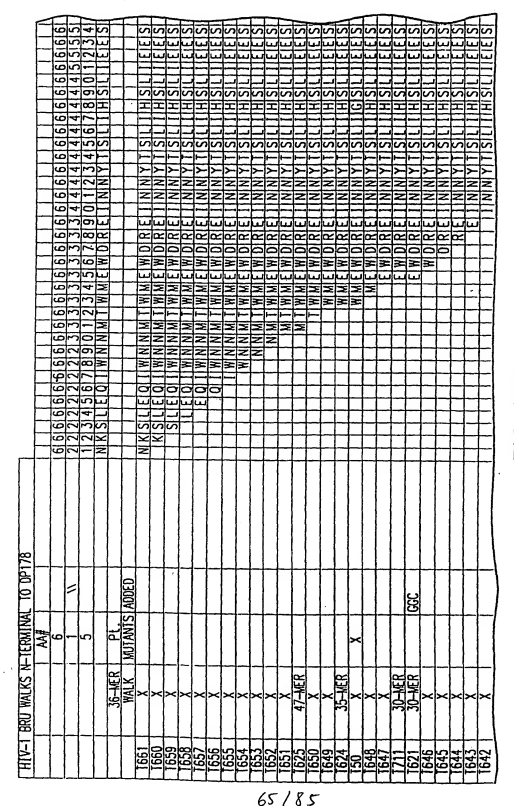
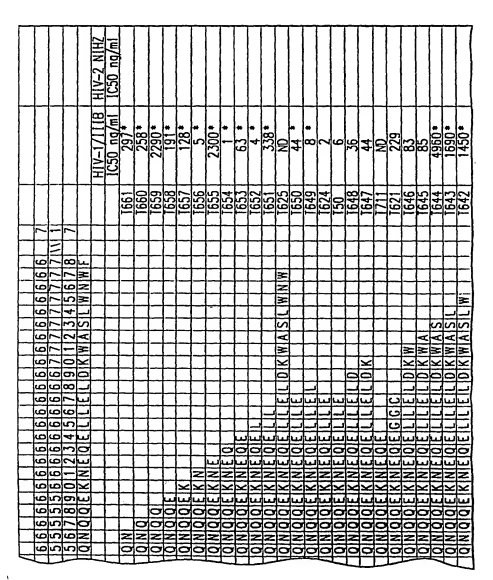


FIG. 49A



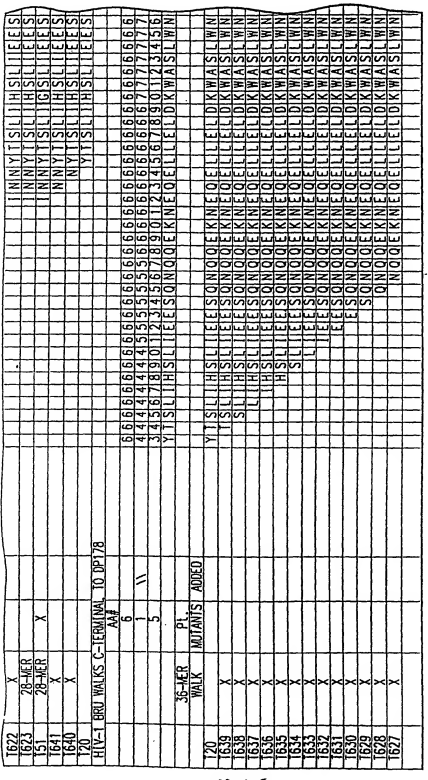


FIG. 49C

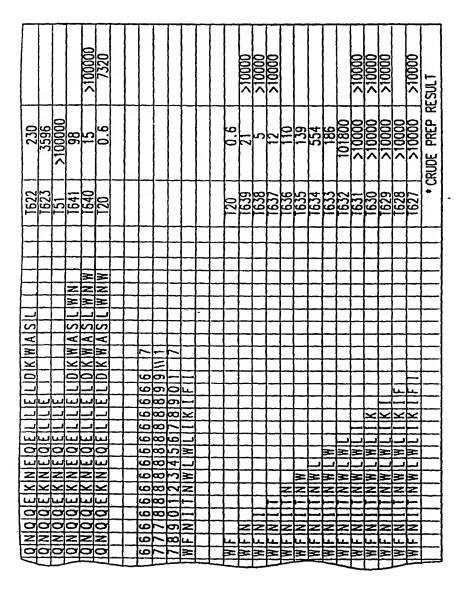


FIG. 49D

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FIG. 49F

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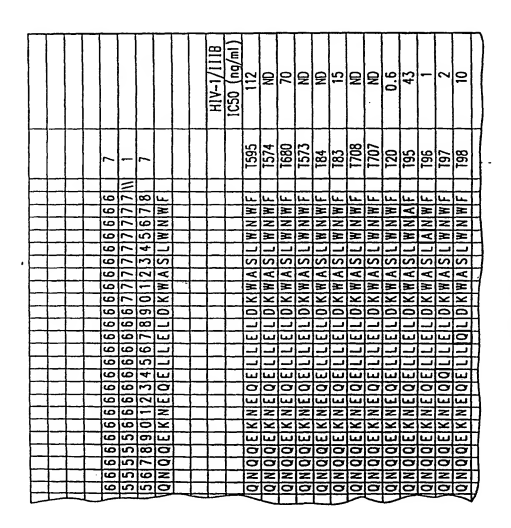
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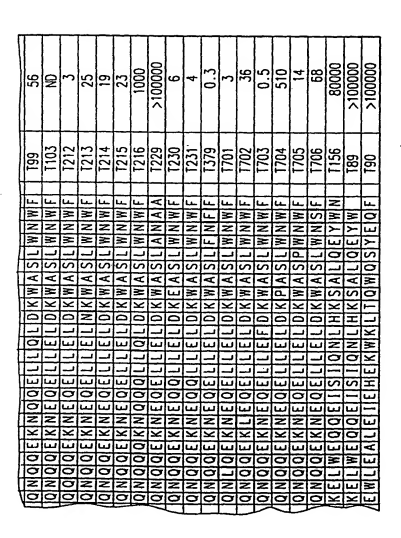
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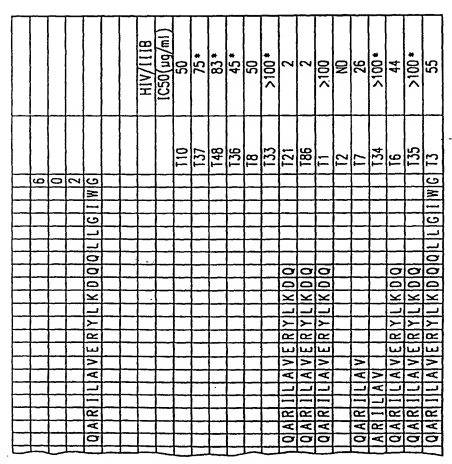
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FIG.51A

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185 A S R K C R A K F K O L L O H Y R E V A A A K K C R A K F K O L L O H Y R E V A A A K K C R A K F K O L L O H Y R E V A A A K K C R A K F K O L L O H Y R E V A A A K K C R A K F K O L L O H Y R E V A A A K K C R A K F K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A A K K K O L L O H Y R E V A A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y R E V A A A A K K K O L L O H Y A A A K K K O L L O H Y A A A K K K O L L O H Y A A A K K K O L L O H Y A A A K K K O L L O H Y A A A K K K O L L O H Y A A A K K K A A A A K K K A A A A A K K K A A A A A K K K A A A A A A A A A A A A A A A A A A A A	Ц				S				S	S	S	က	S	ကျ	S	S			
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FIG.51B

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174 P-L-L-V-L-Q-A-G-F-F-L-L-I-R-1-L-I-I-P-Q-S-L-D-S-W-W-I-S-L-N-F-L-G-G-G-I-I-V-C-L-G-Q-N-S-Q-S-P 220 DOMAIN 1:

F-L-L-1-R-1-L-1-1-1-P-Q-S-L-D-S-W-W-1-S-L-N-F-L-G-G-1-1-V-C-L-G-Q-N-S-Q S-W-D-D-1-1-1-1-1-1-D-D-3-W-M-1-2-T-W-Z-1-1-1-1-1-1-1-W-1-1-1-N-C-1-1-N-C-T-1-N-C-T-1-N-C-T-1-N-S-C-N-W-S-T-N-G-F-F-i(-1-1-R-1-1-1-1-1-P-Q-S-1-D-S-W-W-1-S-1-N-F-1-G-G-1-1-V-C-1-G-Q-N A-G-F-F-L-L-1-R-1-L-1-1-P-Q-S-L-D-S-W-W-1-S-L-N-F-L-G-G-1-1-V-C-L-G-Q Q-A-C-F-F-L-L-1-R-1-L-1-1-P-Q-S-L-D-S-W-W-1-S-L-N-F-L-G-G-1-1-V-C-L-G L-Q-A-G-F-F-L-L-I-R-I-L-I-J-P-Q-S-L-D-S-W-W-I-S-L-N-F-L-G-G-I-I-V-C-L V-L-Q-A-G-F-F-L-L-1-1-1-1-1-0-S-L-D-S-1-N-1-S-L-N-F-L-G-G-1-1-V-C L-V-L-Q-A-G-F-F-L-L-T-R-J-L-T-1-P-Q-S-L-D-S-W-W-T-S-L-N-F-L-G-G-T-T-V [-[-V-L-Q-A-G-F-F-L-L-T-R-]-L-T-P-Q-S-L-D-S-W-W-T-S-L-N-F-L-G-G-T-T

FIG.52A

DOMAIN 11:

223 P-G-Y-R-W-H-C-L-R-R-F-I-I-F-L-F-I-L-L-L-C-L-I-F-L-L-V-L-L-D-Y-Q-G-H-L-P-V-C-P-L-I-P-G-S-S-I-S-I-G-P-C-R-I-C-H-I-591 F-I-I+-1-F-1-F-1-L-1-C-1-F-1-V-1-L-P-Y-Q-G-M-1-P-V-C-P-1-1-G-G-S-S R-F-1-1-F-1-F-1-F-1-L-1-C-1-1-F-1-L-1-V-1-L-1-Y-Q-G-M-1-P-V-C-P-1-1-P-G-S R-R-F-1-1-F-1-F-1-1-1-1-C-1-F-1-1-V-1-1-D-Y-Q-G-14-1-P-V-C-P-1-J-P-G L-R-R-F-1-1-F-L-F-1-L-L-L-C-L-1-F-L-L-V-L-L-D-Y-Q-G-M-L-P-V-C-P-L-1-P C~L~R~R-F-1~1~F~L~F~I~L~L~L~L~L~I~F~L~L~V~L~L~D~Y~Q~G~H~L~P~V~C~P~L~) ╫╫C╌┸┸┰╌╌┇╌┇┼┼┸╌╏╌ **୵-₭-╟╫-ᢗ┤-Ҟ-╀-┊╌┊╌┊╌┊╌┊╌╎╌╎╌╎╌╎╌╌╌╌╌╌╌╌╌╌╌╌╌╌╌** G-Y-R-W-MC-L-R-F-I-I-F-L-F-I-L-L-L-C-L-I-F-L-L-V-L-L-D-Y-Q-G-M-L-P ┢╌╏╌╉╌╫╌╫╌С┰╶R╌╂╌┋╌┇┾╌┸╌┸╌┸╌┸╌┸╌┸╌┸╌┸╌┸╌┸╌┸╌┸╌┸╌┸╌

C-L-]-F-L-V-L-L-D-Y-Q-G-M-L-P-V-C-P-L-]-P-G-S-S-1-S-1-G-P-C-R-1-C-M |-C-|-|-F-|-S-1-S-1-G-W-|-P-V-C-P-|-P-G-S-S-1-S-1-G-P-C-R-1-C |-1-C-|-1-f-|-1-n-1-n-n-n-n-n-n-n-n-1-1-1-n-c-S-2-1-S-1-C-h-1-|-L-L-L-C-L-1+-L-L-L-L-D-Y-Q-G-11-L-P-V-C-P-L-1-P-G-S-S-1-S-1-G-P-C

FIG.52B

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T888	ICSO	Conc. (nM)		6.4	Insoluble	75.6	7.3	28.7		3.5	195	7.2	Insoluble	624	4.8												
Fusion	IC50 µg/ml		0.30	2.6	1.7	3	2.1	1.3	2.1	6.0	0.5	0.5	3.8	1.3	1.6												-
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		, F	7111	1800	1801	T802	T803	T804	T805	T806	T807	T808	T809	T810	T811	T1669	T1670	T1671	T1672	T1673	T1680	T1681	T1682	T1683	T1684		

	Does	Fusion	T83	T1342 FP
<u>n</u>	Substitution		ELAKA (Biot-	-tan-
स	interaction?	1C50	T20)	1071
		(ng/mL		IC 50
		_	1C50	Conc.
			Conc.	(mu)
Ac-YTSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2	Native	2	69	
AC-AAAAIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2		3	1000	
AC-YTSAAASLIEESQNQQEKNEQELLELDKWASLWNWF-NH2	Reduced	37	1000	
ac-ytslihaaaeesqnqqekneqelleldkwaslwnwf-nh2	Inhibited	1757	Neg	
ac-ytslihsliaaaqnqqekneqelleldkwaslwwf-nh2	Native	11	29	
AC-YTSLIHSLIEESAAAQEKNEQELLELDKWASLWNWF-NH2	Inhibited	33	>3000	
AC-YTSLIHSLIEESQNQAAANEQELLELDKWASLWNWF-NH2	Inhibited	545	>3000	
AC-YTSLIHSLIEESQNQQEKAAAELLELDKWASLWNWF-NH2	Inhibited	814	Neg	
AC-YTSLIHSLIEESQNQQEKNEQAAAELDKWASLWNWF-NH2	Native	23	99	
Ac-ytslihslieesqnqqekneqellaaakwaslwwf-nh2	Inhibited	88	1156	
Ac-Ytslihslieesqnqoekneqelleldaaaslwnwf-nh2	Reduced	8	251	
ac-ytslihslieesqnqqekneqelleldkwaaanwf-nh2	Reduced	28	57	
ac-ytslihslieesqnqqekneqelleldkwaslwaa-nh2	Reduced	56	56	
AC-ATSLIHSLIEESQNQQEKNEQELLELDKWASLWNWF-NH2	Native			
Ac-Yaslihslieesqnqqekneqelleldkwaslwnwf-nh2	Native			
Ac-Ytalihslieesonqoekneqelleldkwaslwnwf-nh2	Native			
Ac-Ytsaihslieesqnqqekneqelleldkwaslwnwf-nh2	Reduced			
Ac-ytslahslieesqnqqekneqelleldkwaslwnwf-nh2	Reduced			
ac-ytsliaslieesqnqqekneqelleldkwaslwnwf-nh2	Native			
Ac-Ytslihalieesqnqqekneqelleldkwaslwnwf-nH2	Native			
Ac-Ytslihsaleesqnqqekneqelleldkwaslwnwf-nh2	Reduced			
Ac-YTSLIHSLAEESQNQQEKNEQELLELDKWASLWNWF-NH2	Inhibited			
AC-YTSLIHSLIEESANQQEKNEQELLELDKWASLWNWF-NH2	Native			9
Ac-YTSLIHSLIEESQAQQEKNEQELLELDKWASLWNWF-NH2	Native			
AC-YTSLIHSLIEESQNAQEKNEQELLELDKWASLWNWF-NH2	Inhibited		-	
Ac-Ytslihslieesqnqaekneqelleldkmaslwnwf-nh2	Inhibited			1000
Ac-YTSLIHSLIEESQNQQAKNEQELLELDKWASLWNWF-NH2	Native			14
Ac-YTSLIHSLIEESQNQQEANEQELLELDKWASLWNWF-NHZ	Native			40
AC-YTSLIHSLIEESQNQQEKAEQELLELDKWASLWNWF-NH2	Inhibited			>1000
Ac-YTSLIHSLIEESQNQQEKNAQELLELDKWASLWNWF-NH2	Reduced			200
Ac-YTSLIHSLIEESQNQQEKNEAELLELDKWASLWNWF-NH2	Native			27
AC-YTSLIHSLIEESQNQQEKNEQELLALDKWASLWNWF-NH2	Native			27
Ac-Ytslihslieesqnqqekneqelleadkwaslwnwf-nh2	Reduced			250
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Fig. 54

Internional application No. PCT/US00/35727

IPC(7) : US CL :	SSIFICATION OF SUBJECT MATTER C12Q 1/70; G01N 33/53; A61K 38/00, 39/21 435/5, 7.1; 530/300; 424/188.1, 208.1		
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C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.
Y	CHEN, CH., et al., A Molecu Immunodeficiency Virus (HIV) Type 1 Anti-HIV Activity of gp41 Derivatives: J. Virol. June 1995. Vol. 69. No. 6. document.	TM Protein Determines the Implication for Viral Fusion.	1-3
Y	US 5,656,480 A (WILD et al.) 1 document.	2 August 1997, see entire	1-3
Y	US 5,464,933 A (BOLOGNESI et al entire document.		1-3
Furth	ner documents are listed in the continuation of Box C	See patent family annex.	
	ecial categories of cited documents:	"I" later document published after the inte date and not in conflict with the appl	lication but cited to understand
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